This Page Is Inserted by IFW Operations and is not a part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

IMAGES ARE BEST AVAILABLE COPY.

As rescanning documents will not correct images, please do not report the images to the Image Problem Mailbox.



PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau

INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:

C07D 471/08. 453/02 // (C07D 487/08. 209:00, 209:00)

(11) International Publication Number:

140 95/03306

- C07D 487/08, 453/06, A01N 43/90, 43/82,
- (43) International Publication Date:

2 February 1995 (02.02.95)

(21) International Application Number:

PCT/US94/08+04

(22) International Filing Date:

21 July 1994 (21.07.94)

- (74) Agents: COSTELLO, James, A. et al.; E.I. du Pont de Nemours and Company, Legal/Patent Records Center, 1007 Market Street, Wilmington, DE 19898 (US).
- (30) Priority Data: 22 July 1993 (22.07.93) 08/095.876 22 November 1993 (22.11.93) 08/156.197
 - US US

- (81) Designated States: AM, AU, BB, BG, BR, BY, CA, CN, CZ, FL GE, HU, IP, KG, KP, KR, KZ, LK, LT, LV, MD, MG, MN. NO. NZ, PL. RO. RU. SL SK, TJ, TT, UA, US, UZ, VN, European patent (AT. BE. CH. DE. DK. ES. FR. GB. GR. E. IT. LU. MC. NL. PT. SE). OAPI patent (BF. BJ. CF, CG, CI, CNL GA, GN, ML, MR, NE, SN, TD, TG). ARIPO patent (KE, MW, SD).

(60) Parent Applications or Grants (63) Related by Continuation

US Filed on

Filed on

US

08/095.876 (CIP) 22 July 1993 (22.07.93) 22 November 1993 (22.11.93)

08/156.197 (CIP)

Published

With international search report.

- (71) Applicant (for all designated States except US): E.I. DU PONT DE NEMOURS AND COMPANY [US/US]: 1007 Market Street Wilmington, DE 19898 (US).
- . (72) Inventor; and
- (75) Inventor/Applicant (for US only): PIOTROWSKI, David, Walter [US/US]: 1712 Christiana Meadows, Bear, DE 19701 (US).
- (54) Title: ARTHROPODICIDAL AZACYCLIC HETEROCYCLES

(11)

(1)

€-2

₹.5

Ç-:

(57) Abstract

Arthropodicidal compounds having formula (I) and compositions comprising compounds of formula (II) and use of said compounds to control arthropods, wherein Q is selected from the group Q-1, Q-2, Q-3 and Q-4; and Q¹ is selected from the group Q-5, Q-6 and Q-7.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pumphiets publishing international applications under the PCT.

21 9200		-		MR	Mariusu
AT	Austra	C.B	Cared Kingdom	70W	Malast
	Yesten	GΕ	Çest\$:±	NE	Niger
AU 83	Buthake	GN	Cuines	NL.	אכטטישטא א
	Seignen	GR	Great	80	Namely
BE	Surtan Fue	នប	Humg sty	NO NZ	New Testing
BF	Brigger	Œ	بحديها		Poluad
8G	-	ιť	انتخ	PL	9 ಗಿದ್ದಾಗ
3.1	Beaus .	JP.	ligina	ट्रा	-
88	8:교	KE	Keep s	80	Russus Federates
BY.	Selara	КG	Kirrorus	RĽ	
ÇA	Curata	KP	Democratic Profile a Reportise	SD	حكور
CF	Ceers 4 ಗಗರಾ ನಿರ್ಗಾಸಿ	•••	el Korea	38	Swider
CG	Cocks	KR	Republic of Korea	51	Slovesta
CII	المات المات	XZ	Kumasa	SK	Slavakia
Cl	Circ dilinose	ü	Lindrakes	\$N	Secretal
CM	Callering	Ľ.	Sn Leaka	מז	C ¹
C.	Com	LÜ	lames	rc	Togo
C2	Carrier atu	LV	Lania	LT	Tajdisus
CZ	Circa Republic	410	Micara	π	Traded Lot Tradego
DΕ	German's	אנט	Republic of Moldova	Ľ A	ارتبته
DΚ	Central			t's	لاميحة لامنح والكحجمة
F2.	ت ج دنه ج	MG	Matugustat Matugustat	ĽZ	ويبونيعت
F١	لحدادع	\fL		VN	Vid Nia
FR	France .	,0,	Morphia		
G.V	ರೆಚುಂತ				

10.

ı

ARTHROPODICIDAL AZACYCLIC HETEROCYCLES

The present invention comprises compounds useful for the control of arthropods. EP 412.798 A2 discloses pyridinyl-substituted azabicyclic compounds for use in dementia provided the pyridinyl ring is not attached to the 2-carbon position of the azabicyclic moiety. J. Gen. Chem. U.S.S.R., (1963), 33, 3345 discloses 2-(3-pyridinyl)-1-azabicyclo[2.2.2]octane, however, no utility is taught. WO 93/14636 discloses azacyclic rings substituted with an optionally substituted oxadiazolyl or thiadiazolyl ring as insecticides. None of these references specifically teaches the compounds of the present invention.

SUMMARY OF THE INVENTION

This invention pertains to compounds of Formula I, including all geometric and stereoisomers and agriculturally suitable salts thereof. The compounds are:

$$\mathbb{R}^1$$
 \mathbb{R}^2

15 wherein:

O is selected from the group

where the broken line represents an optional chemical bond:

20 R¹ and R² are independently selected from the group H, halogen, C₁-C₆ alkyl, C₁-C₅ haloalkyl, C₂-C₆ alkenyl, C₂-C₆ haloalkenyl, C₂-C₆ alkynyl, C₂-C₆ haloalkynyl, C₃-C₆ eyeloalkyl, C₃-C₆ halocycloalkyl, CN, SCN, NO₂, N(R⁵/R⁶, OR⁵, C(O)R⁵, C(O)OR⁵, C(O)N(R⁵)R⁶, SR⁵, S(O)R⁵, S(O)₂R⁵, S(O -N(R⁵)R⁶) and C₁-C₆ alkyl substituted with 1 or 2 groups

PCT/US94/08404

2

5	independently selected from NO ₂ , CN, C ₁ -C ₃ alkylthio, C ₁ -C ₃ alkoxy, C ₁ -C ₃ haloalkoxy, C ₂ -C ₄ alkylcarbonyl and C ₂ -C ₄ alkoxycarbonyl; R ² being attached to any unsubstituted aromatic ring carbon; and R ¹ and R ² are not both hydrogen when Q is Q-1 or Q-4, n is 1, R ³ is H and q is 2; R ³ , which is attached to any carbon of the azabicyclic ring including the carbon directly attached to the heterocyclic aromatic ring, is selected from the group H, halogen, C ₁ -C ₆ alkyl, C ₁ -C ₆ haloalkyl, C ₂ -C ₆ alkenyl, C ₂ -C ₆ haloalkenyl, C ₂ -C ₆ alkynyl, C ₂ -C ₆ haloalkynyl, C ₃ -C ₆ cycloalkyl, C ₃ -C ₆ halocycloalkyl, CN, SCN, NO ₂ , N(R ⁷)R ⁸ , OR ⁷ , C(O)R ⁷ , C(O)OR ⁷ , C(O)OR ⁷ , C(O)OR ⁷ , S(O) ₂ R ⁸ and C ₁ -C ₆ alkyl
	substituted with 1 or 2 groups independently selected from the group NO ₂ . CN, C ₁ -C ₃ alkytthio, C ₁ -C ₃ alkoxy, C ₁ -C ₃ haloalköxy, C ₂ -C ₄ alkylearbonyl and C ₂ -C ₄ alkoxycarbonyl; R ² is selected from the group H, C ₁ -C ₆ alkyl, C ₃ -C ₆ cycloalkyl, C ₁ -C ₆ haloalkyl,
15	C ₂ -C ₆ alkenyl, C ₂ -C ₆ alkynyl, N(R ⁹)R ¹⁰ , C(O)R ⁹ , C(O)OR ⁹ , C(O)OR ⁹ , C(O)N(R ⁹)R ¹⁰ , SR ⁹ , S(O)R ⁹ , S(O) ₂ R ⁹ · S(O) ₂ N(R ⁹)R ¹⁰ , benzyl and CH(CH ₃)Ph; provided when any of R ¹ , R ² , R ³ or R ⁴ is S(O)R ⁵ , S(O) ₂ R ⁵ , S(O)R ⁷ , S(O) ₂ R ⁷ , S(O)R ⁹ , or S(O) ₂ R ⁹ then R ⁵ , R ⁷ and R ⁹ are other than H; R ⁵ , R ⁶ , R ⁷ , R ⁸ , R ⁹ and R ¹⁰ are independently selected the group H, C ₁ -C ₆ alkyl,
20	C ₁ -C ₆ haloalkyl, C ₃ -C ₆ cycloalkyl, phenyl optionally substituted with 1 or 2 substituents independently selected from W, and benzyl optionally substituted with 1 or 2 substitutents independently selected from W; W is selected from the group halogen, NO ₂ , CN, C ₁ -C ₃ alkyl, C ₁ -C ₃ haloalkyl, C ₁ -C ₃ alkylthio, C ₁ -C ₃ alkoxy, C ₁ -C ₃ haloalkoxy, C ₂ -C ₄ alkylcarbonyl and
25	C ₂ -C ₄ alkoxycarbonyl;
	m and n are independently 0, 1 or 2;
	p is 1 or 2; and
	q is 1, 2 or 3.
30	Preferred Compounds A are compounds of Formula I wherein Q is Q-1.
	Preferred Compounds B are compounds of Preferred A wherein:
	R^{T} is selected from the group H, halogen and C_{T} - C_{2} alkyl:
	R ² is selected from the group H and CI;
35	R3 is selected from the group H, halogen, C ₁ -C ₆ alkyl and OR7;

 R^5 is selected from the group H and $C_1\text{-}C_4$ alkyl ; and

n is Car I.

is:

3

Specifically preferred for biological activity is Compound C of Preferred B which

7-(6-chloro-3-pyridinyl)-1-azabicyclo[2.2.1]heptane.

This invention also includes arthropodicidal compositions and method of use in both agronomic and nonagronomic environments wherein the arthropodicidal compounds are those of the formula:

10

wherein:

Q1 is selected from the group

$$(R^{3})_{p} = (CH_{2})_{n}$$

$$(CH_{2})_{p} = (CH_{2})_{m}$$

$$(CH_{2})_{q} = and$$

$$(CH_{2})_{m} = (CH_{2})_{m}$$

$$(R^{3})_{p} = (CH_{2})_{m} = (CH_{2})_{m}$$

$$(R^{3})_{p} = (CH_{2})_{m} = (CH_{2})_{m$$

15

where the broken line represents an optional chemical bond;

V. X. Y and Z of the ring are each independently selected from the group O, S, N.

-C(R¹)-, -C(R¹)=C(R²)-, -C(R¹)=N- and -N(R¹¹)-; provided that (i) no more than one of V, X, Y or Z is -C(R¹)=C(R²)-, -C(R¹)=N-, -N(R¹¹)-, O

or S. (ii) at least one of V, X, Y or Z is N, (iii) when the ring is a 5-membered ring containing two N and one O or S, and Q is Q-6, then the ring is attached to the 2-position of Q¹ and (iv) when the ring is a 5-membered ring containing two N and one O or S, then Q is other than Q-5;

R¹ and R² are independently selected from the group H, halogen, C₁-C₆ alkyl.

C₁-C₆ haloalkyl, C₂-C₆ alkenyl, C₂-C₆ haloalkenyl, C₂-C₆ alkynyl, C₂-C₆ haloalkynyl, C₃-C₆ cycloalkyl, C₃-C₆ haloayeloalkyl, CN, SCN, NO₂, N(R⁵)R⁶, OR⁵, C(O)R⁵, C(O)OR⁵, C₁O)N(R⁵)R⁶, SR⁵, S(O)R⁵, S(O)₂R⁵, S(O)₂R⁵, S(O)₂N(R⁵)R⁶ and C₁-C₆ alkyl substituted with 1 or 2 groups independently selected from the group NO₂, CN, C₁-C₃ alkylthio, C₁-C₃

alkoxy, C₁-C₂ haloalkoxy, C₂-C₄ alkylearbonyl and C₂-C₄ alkoxycarbonyl;

4

Re, which is attached to any earbon of the azacyclic ring including the carbon
directly attached to the heterocyclic aromatic ring, is selected from the
group H. halogen, C1-C6 alkyl, C1-C6 haloalkyl, C2-C6 alkenyl, C2-C6
haloalkenyl, C2-C6 alkynyl, C2-C6 haloalkynyl, C3-C6 cycloalkyl, C3-C6
halocycloalkyl, CN, SCN, NO ₂ , $N(R^7)R^8$, OR^7 , $C(O)R^7$, $C(O)OR^7$,
$C(O)N(R^7)R^8$, SR^7 , $S(O)R^7$, $S(O)_2R^7$, $S(O)_2N(R^7)R^8$ and C_1 - C_6 alkyl
substituted with 1 or 2 groups independently selected from NO2, CN,
C_1 - C_3 alkylthio, C_1 - C_3 alkoxy, C_1 - C_3 haloalkoxy, C_2 - C_4 alkylcarbonyl and
C ₂ -C ₄ alkoxycarbonyl;
$ m R^4$ and $ m R^{11}$ are independently selected from the group H, $ m C_1$ - $ m C_6$ alkyl, $ m C_3$ - $ m C_6$
cycloalkyl, C ₁ -C ₆ haloalkyl, C ₂ -C ₆ alkenyl, C ₂ -C ₆ alkynyl, N(R ⁹)R ¹⁰ ,
$C(O)R^9$, $C(O)OR^9$, $C(O)N(R^9)R^{10}$, SR^9 , $S(O)R^9$, $S(O)_2R^9$.
$S(O)_2N(R^9)R^{10}$, benzyl and $CH(CH_3)Ph$; provided when any of R^1 , R^2 , R^3
or \mathbb{R}^4 is $S(O)\mathbb{R}^5$, $S(O)_2\mathbb{R}^5$, $S(O)\mathbb{R}^7$, $S(O)_2\mathbb{R}^7$, $S(O)\mathbb{R}^9$, or $S(O)_2\mathbb{R}^9$ then \mathbb{R}^5 .
R ⁷ and R ⁹ are other than H;
R5, R6, R7,R3, R9 and R10 are independently selected the group H, C1-C6 alkyl.
C ₁ -C ₆ haloalkyl, C ₃ -C ₆ cycloalkyl, phenyl optionally substituted with 1 or 2
substituents independently selected from W, and benzyl optionally
substituted with 1 or 2 substitutents independently selected from W;
W is selected from the group halogen, NO ₂ , CN, C ₁ -C ₃ alkyl, C ₁ -C ₃ haloalkyl,
C_1 - C_3 alkylthio, C_1 - C_3 alkoxy, C_1 - C_3 haloalkoxy, C_2 - C_4 alkylcarbonyl and
C ₂ -C ₄ alkoxycarbonyl;
m and n are independently 0, 1 or 2;
p is 1 or 2; and
q-is-1 , 2 -or-3.

Preferred Method A is the method wherein:

 R^{1} is selected from the group H, halogen and $C_{1}\text{-}C_{2}$ alkyl;

R2 is selected from the group H and CI;

 R^3 is selected from the group H, halogen, C_1 - C_6 alkyl and OR^7 ;

 R^5 is selected from the group H and C_1 - C_4 alkyl; and

m and n are 0 or 1.

Preferred Method B is the method of Preferred A wherein:

V is N: X is -C(R¹)=C(R²)-; Y and Z are -C(R¹)-; and Q¹ is Q-6.

20

30

35

5

Preferred Method C is the method of Preferred A wherein:

V is N;

X is S:

Y and Z are -C(R1)-; and

Q! is Q-6.

Preferred Method D is the method of Preferred A wherein: the ring contains two N and one O or S; and O1 is Q-6.

Compounds of this invention can exist as one or more stereoisomers. The various stereoisomers include enantiomers, diastereomers and geometric isomers. One skilled in the art will appreciate that one stereoisomer may be more active than the others and how to separate stereoisomers. Accordingly, the present invention comprises racemic and optically active compound(s) of Formulae I and II as well as agriculturally suitable salts thereof. The term optically active compound(s) includes individual stereoisomers, mixtures of stereoisomers enriched in one stereoisomer, and optically active mixtures of compounds.

In the above recitations, the term "alkyl", used either alone or in compound words such as "alkylthio" or "haloalkyl" denotes straight-chain or branched alkyl, such as, methyl, ethyl, n-propyl, i-propyl, or the different butyl, pentyl or hexyl isomers. "Alkenyl" denotes straight-chain or branched alkenes such as ethenyl, 1-propenyl, 2-propenyl, and the different butenyl, pentenyl and hexenyl isomers. "Alkenyl" also denotes polyenes such as 1.3-hexadiene. "Alkynyl" denotes straight-chain or branched alkynes such as ethynyl, 1-propynyl, 3-propynyl and the different butynyt, pentynyl and hexynyl-isomers. "Alkynyl"-can-also-denote-moieties-comprised of multiple triple bonds such as 2,4-hexadiyne. "Alkoxy" denotes methoxy, ethoxy, n-propyloxy and isopropyloxy isomers. "Cycloalkyl" denotes cyclopropyl, cyclobutyl, cyclopentyl and cyclohexyl. The term "halogen", either alone or in compound words such as "haloa!kyl", denotes fluorine, chlorine, bromine or iodine. Further, when used in compound words such as "haloalkyi", said alkyl may be partially or fully substituted with halogen atoms which may be the same or different. Examples of "haloalkyl" include F3C, CiCH2, CF3CH2 and CF3CCl2. Examples of "haloalkenyl" include (CI)2C=CHCH2 and CF3CH2CH=CHCH2. Examples of "haloalkynyl" include HC = CCHCI, $CF_3C = C$, $CCI_3C = C$ and $FCH_2C = CCH_2$. The total number of carbon atoms in a substituent group is indicated by the " C_i - C_i " prefix where i and j are numbers from 1 to 6. For example, C_1 - C_3 alkyl designates methyl through propyl; C_2 alkoxy designates CH₂CH₂O; C₃ alkoxy designates CH₃CH₂CH₂O or (CH₃)₂CHO.

10

15

20

ó

Examples of "alkoxycarbonyl" include $CH_3OC(=0)$, $CH_3CH_2OC(=0)$, $CH_3CH_2CH_2OC(=0)$ and $(CH_3)_3CHOC(=0)$.

Examples of Formula II heterocyclic rings, exclusive of Q¹, include pyrrolyl, imidazolyl, pyrazolyl, oxazolyl, thiazolyl, isoxazolyl, isothiazolyl, oxadiazolyl, thiadiazolyl, triazolyl, pyridyl, pyridazinyl, pyrimidinyl, pyrazinyl, oxazinyl, thiazinyl, oxadiazinyl, thiadiazinyl, triazinyl and tetrazinyl.

The 2-position of the azabicyclic ring is defined as any carbon atom directly attached to the nitrogen atom of the azabicyclic ring. The 3-position is defined as any carbon atom directly attached to any carbon atom which is directly attached to the nitrogen atom of the azabicyclic ring.

When a compound is substituted with a substituent bearing a subscript that indicates the number of said substituents can exceed 1, said substituents (when they exceed 1) are independently selected from the group of defined substituents. Similarly, when a compound is substituted with a substituent which occurs more than once, said substituent is independently selected from the group defined for said substituent.

DETAILS OF THE INVENTION

Compounds of Formula I (Q-1) and Formula II (Q-6) can be prepared by intramolecular alkylation of amino compounds of Formula III where G is a suitable leaving group such as a halogen or sulfonate. The reaction can be performed in the absence or presence of a base in a suitable solvent. Suitable solvents include tetrahydrofuran, diethyl ether and dimethoxyethane. Examples of typical bases include sedium methoxide, potassium t-butoxide and sodium hydride. The reactions can be run at temperatures in the range of 25°C to reflux. This method is similar to procedures described in the art (Chimia (1976) 30, 60). Scheme I illustrates this reaction.

25

30

Schene I

$$(CH_{2})_{n}$$

$$(CH_{2})_{q}$$

$$(CH_{2})_{q}$$

$$(CH_{2})_{q}$$

$$(R^{3})_{p}$$

Compounds of Formula III can be prepared by reaction of the hydroxy compounds of Formula IV with a halogenating agent such as thionyl chloride, phosphorous tribromide or a sulfonylating agent such as methanesulfonyl chloride or *p*-toluenesulfonyl chloride using procedures that are known to one skilled in the art

7

(March, J. Advanced Organic Chemistry, John Wiley & Sons, New York, 3rd ed. (1985) 1151). Scheme II illustrates this transformation.

Scheme II

$$\begin{array}{c} \text{OH} \\ \text{Z-V} \\ \text{CH}_{2} \text{I}_{n} \\ \text{CH}_{2} \text{I}_{q} \end{array}$$

IV (where $R^{\lambda} = H$)

III (where $R^X = H$)

G is halogen or sultonate

5

10

15

Formula IV compounds can be prepared by reduction of ketones with a reducing agent such as sodium borohydride, borane, and lithium aluminum hydride using procedures that are known to one skilled in the art (March, J. Advanced Organic Chemistry, John Wiley & Sons, New York, 3rd ed. (1985) 1147). Scheme III illustrates this transformation.

Scheme III

OH

$$Z=V$$
 $(CH_2)_n$
 $(CH_2)_q$
 $(CH_2)_q$

Alternatively. Formula IV compounds can be prepared by addition of an appropriate organometallic compound VI (where M = Li, Mg, Cu, Zn) to a carbonyl compound VII in a suitable solvent such as diethyl ether, tetrahydrofuran and dimethoxyethane. The reaction can be run at temperatures from -78°C to 35°C. Scheme IV illustrates this transformation.

10-

S

$$\bigcup_{Z} \bigcup_{Q} R$$

$$\forall H \quad (R^Z = H)$$

(where RX + H)

Compounds of Formula V can be prepared by deprotection of compounds of Formula VIII using procedures known to one skilled in the art (Greene, Protective Groups in Organic Synthesis, 2nd ed. (1991) 315-348). Scheme V illustrates this 5 transformation.

Scheme V

$$(CH_2)_n \qquad X - Y$$

$$(CH_2)_n \qquad (CH_2)_n \qquad (CH_2)_n$$

$$(CH_2)_q \qquad (CH_2)_q \qquad (CH_2)_q$$

$$(CH_2)_q \qquad (CH_2)_q \qquad (CH_2)_q$$

$$(CH_2)_q \qquad (CH_2)_q \qquad (CH_2)_q$$

$$(CH_2)_q \qquad (CH_2)_q \qquad (CH_2)_q \qquad (CH_2)_q \qquad (CH_2)_q$$

VIII (Ry is a protecting group)

Compounds of Formula VIII can be prepared by de-alkylation of Formula IX compounds with a chloroformate derivative such as 1-chloroethyl chloroformate, benzyl chloroformate, and 2.2.2-trichloroethyl chloroformate in a suitable solvent such as methylene chloride, chloroform, and 1.2-dichloroethane. The reactions are usually run at temperatures in the range of 25°C to the reflux temperature of the particular 15 solvent. Scheme V! illustrates this transformation.

Scheme VI

Scheme VI

$$(CH_2)_n \qquad (CH_2)_n \qquad (CH_2)_n \qquad (CH_2)_q$$

$$(CH_2)_q \qquad (CH_2)_q \qquad (CH_2)_q$$

$$(CH_2)_q \qquad (CH_2)_q \qquad (CH_2)_q$$

Compounds of Formula IX can be prepared by addition of an organometallic compound of Formula VI to carbonyl compounds of Formula VII in an analogous procedure described for Formula IV compounds. Scheme VII illustrates this transformation.

Scheme VII $M = (where R^{\prime} = H)$ Vil (where RZ = Cl. N (OMe) Me. ÍΧ alkoxy)

Carbonyl compounds of Formula VII where Rz is -N(OMe)Me can be prepared from carboxylic acid derivatives using procedures known in the art (Org. Prep. Proc. (1993) 25, 15%

Organometallic compounds of Formula VI can be prepared by reaction of Formula X with a reactive metal species, such as magnesium, using procedures similar to those described in the art (J. Med. Chem. (1965) 8, \$29). One skilled in the art will recognize this as a Grignard reagent. Alternatively, Formula VI compounds (M=Li) can be obtained by lithium halogen exchange using procedures known to one skilled in the art (March, J. Advanced Organic Chemistry, John Wiley & Sons, New York, 3rd ed. (1985) 1169 .

15

Schene VIII

Scheme VIII

$$(CH_{2})_{\pi}$$

$$(CH_{2})_{\eta}$$

$$(CH_{2})_{\eta}$$

$$(CH_{2})_{\eta}$$

$$(CH_{2})_{\eta}$$

$$(CH_{2})_{\eta}$$

$$(CH_{2})_{\eta}$$

X (G=halogen)

(M is magnesiumor lithium reagent)

M (M is Mg or Li)

Alternatively, compounds of Formula I (Q-1) and Formula II (Q-6) can be prepared by depretection of salts of Formula XI by hydrogenation in the presence of a suitable catalyst and solvent. Suitable catalysts include palladium on carbon and platinum oxide. Appropriate solvents are methanol, ethanol and ethyl acetate. Similar procedures are described in the art (J. Chem. Soc., Perkin Trans. 1, (1991) 1091). Scheme IX illustrates this transformation.

Scheme IX

$$(CH_2)_n$$

$$(R^3)_p$$

$$(R^$$

10

15

Salts of Formula XI can be prepared by cyclization of Formula III compounds in a suitable solvent. Suitable solvents are ethanol, 1,2-dichloroethane and toluene. The reaction can be run at temperatures in the range of 0°C to reflux. Scheme X illustrates this reaction.

Schene X

$$CH_{2^{1}n}$$
 $X = Y$
 $CH_{2^{1}n}$
 $X = Y$
 $CH_{2^{1}n}$
 $CH_{2^$

III (where R = C₁-C₆ alky), benzy).
Gis helogen or sultonate)

ΧI

15

One skilled in the art will recognize that one substituent can be converted into another. For example, compound I ($V=CR^T$, $R^T=CI$) can be converted into I ($V=CR^T$, $R^T=OMe$) by displacement with methoxide, or converted into I ($V=CR^T$, $R^T=alkyI$) by reaction with an organostannane under palladium catalysis.

5 Scheme XI illustrates this transformation.

Scheme XI

$$(CH_2)_n$$

$$(CH_2)_q$$

$$(R^3)_p$$

$$(R^3)_p$$

I (where V = CR1, R1 = halogen)

I (where V = CR¹, R¹ = alkyl, alkoxy, thioalkyl, alkenyl, etc.)

Compounds of Formula I (Q-2) and Formula II (Q-7) can be prepared by cycloaddition of compounds of Formula XII with imines of Formula XIII. The reaction can be performed in the absence or presence of an acid such as zinc chloride, boron trifluoride, and hydrogen chloride. Suitable solvents are dichloromethane, toluene, tetrahydrofuran, and water. The reactions can be run at temperatures ranging from -78°C to reflux temperature of the solvent. It will be recognized by those skilled in the art that fully saturated azabicyclic analogs can be prepared by simple reduction of the olefia. Scheme XII illustrates this transformation.

Scheme XII

L(Q-2) and H (Q-7)

Alternatively, compounds of Formula I (Q-2) and Formula II (Q-7) can be prepared by intramolecular alkylation of Formula XIV compounds in a similar fashion to methods described for Formula I (Q-1) compounds. Scheme XIII illustrates this transformation.

5

Scheme XIII

$$(CH_2)_n$$

$$(CH_2)_m$$

$$(CH_2)_$$

G is halogen or sulfonate

I (Q-2) and ∏ (Q-7)

Imines of Formula XIII can be prepared by condensation of Formula VII carbonyl compounds with a primary amine (R⁴NH₂) using procedures known to one skilled in the art (March, J. Advanced Organic Chemistry; John Wiley & Sons, New York, 3rd ed. (1985) 465). Scheme XIV illustrates this transformation.

Scheme XIV

Y=
$$\frac{X}{Z}$$
 + R^4NH_2 $\frac{Y-X}{Z}$ R^2

VII. (where $R^2 = H$, alkyl)

15

10

Formula I (Q-3) and Formula-II-(Q-7) compounds can be prepared by cycloaddition of compound XV using procedures known to one skilled in the art (J. Am. Chem. Soc. (1985) 107, 1768). It will be recognized by one skilled in the art that fully saturated azabicyclic analogs can be prepared by simple reduction of the olefin by known methods. Scheme XV illustrates this transformation.

BEST AVAILABLE COPY

13 Scheme XV

$$(CH_{2})_{n}$$

$$(CH_{2})_{n}$$

$$(CH_{2})_{n}$$

$$(CH_{2})_{n}$$

$$(R^{3})_{p}$$

$$(R^$$

Formula XV compounds can be prepared by reacting heterocycles of
Formula XVI with olefins XVII under palladium catalysis in the presence of base.
Typical solvents include acetonitrile, dimethylformamide, and tetrahydrofuran. Typical bases include triethylamine, disopropylethylamine and sodium bicarbonate. Reaction temperatures range from room temperature to the reflux temperature of the particular solvent. Scheme XVI illustrates this transformation.

Schene XVI

$$C = C. Br. I. OTi$$

Schene XVI

$$(R^{3})_{p}$$

$$(CH_{2})_{n}$$

$$Z = C. Br. I. OTi$$

NV

Alternatively, compounds of Formula I (Q-3) and Formula II (Q-7) can be prepared by intramolecular alkylation of Formula XVIII compounds by procedures

described for Formula I (Q-1) compounds. Scheme XVII illustrates this transformation.

G is halogen or sultonate

5

Formula XVIII compounds can be prepared by addition of organometallic compounds of Formula XIX to imines XX. Typical solvents can include tetrahydrofuran and diethylether. Typical reaction temperatures range from -78°C to room temperature. Scheme XVIII illustrates this transformation.

10

Scheme XVIII

$$(R^3)_p$$
 $(CH_2)_m$
 $(R^3)_p$
 $(CH_2)_n$
 (CH_2)

protected hydroxyl

NAM

Imines of Formula XX can be prepared by condensation of a primary amine R4NH2 with ketones XXI under conditions known to one skilled in the art (March, J. Advanced Organic Chemistry; John Wiley & Sons, New York, 3rd ed. (1985) 465). Scheme XIX illustrates this transformation.

protected hydroxyl

Scheme XIX

$$(CH_2)_{m}$$

$$(CH_2)_{n}$$

$$(CH_$$

Organometallic compounds of Formula XIX can be prepared by methods known in the art see, for example, EP 492,902-A1.

Compounds of Formula II (Q-6) can be synthesized by addition of organometallic compounds XIX to azabicyclic ketones XXII followed by chlorination, elimination, and hydrogenation as described in the art. See, for example, EP 412,798. Scheme XX depicts this transformation.

Scheme XX

$$(R^{3})_{p} \xrightarrow{(CH_{2})_{q}} (CH_{2})_{q} \xrightarrow{(CH_{2})_{q}} (R^{3})_{p} \xrightarrow{(CH_{2})_{n}} (CH_{2})_{n} \xrightarrow{($$

Compounds of Formula II (Q-6) can be synthesized from azabicyclic esters XXIII and compounds XXIV, in a manner analogous to procedures described in the art. See, for example, EP 323,864. Scheme XXI depicts this transformation.

Schene XXI

$$(R^3)_p \xrightarrow{CH_2)_q} OH \longrightarrow (R^3)_p \xrightarrow{(CH_2)_q} (CH_2)_q$$

$$(CH_2)_q \xrightarrow{(CH_2)_q} (CH_2)_q$$

$$(CH_2$$

5

10

Compounds of Formula II (Q-7) can be prepared from compounds of Formula XXV by procedures described for Formula II (Q-6) compounds. Scheme XXII depicts this transformation.

Scheme XXII

OH

$$CO_2R^a$$
 R^b
 R^b
 $R^a = C_1 \cdot C_6 alkyl$
 XXV
 XXV
 CO_2R^a
 R^b
 R^b

Compounds of Formula XXV can be prepared by hydrogenation of Formula XXVI compounds, which are described in the art (*J. Chem. Soc., Perkin Trans. I* (1991), 1337). Scheme XXIII depicts this transformation.

10

5

Scheme XXIII

$$(CH_2)_q$$

$$CO_2R^a$$

$$R^a = C_1 \cdot C_6 \text{ alky} 1$$

$$XXV$$

Compounds of Formula II (Q-5) can be synthesized by addition of organometallic compounds XIX to azacyclic ketones XXVI, followed by chlorination and elimination.

See, for example, J. Med. Chem. (1992) 35, 4011. Scheme XXIV depicts this transformation.

Schene XXIV

$$(R^{3})_{p}$$

15

25

17

Compounds of Formula I (Q-4) and Formula II (Q-6) can be synthesized by reacting a heterocycle of Formula XVI with an olefin XXVII under palladium catalysis in the presence of base. Typical solvents include acetonitrile, dimethylformamide, tetrahydrofuran, and dimethylsulfoxide. Reaction temperatures range from room temperature to the reflux temperature of the particular solvent. Scheme XXV illustrates this transformation.

Scheme XXV

G

$$CH_{2}^{1}_{n}$$
 $CH_{2}^{1}_{n}$
 $CH_{2}^{1}_{n}$

Olefins of the Formula XXVII can be prepared by using procedures known in the art (Helv. Chim. Acta. (1957) 40, 2170).

It is recognized that some reagents and reaction conditions described above for preparing compounds of Formulae I and II may not be compatible with certain functionalities present in the intermediates. In these instances, the incorporation of protection/deprotection sequences into the synthesis will aid in obtaining the desired products. The use and choice of appropriate protecting groups will be apparent to one skilled in chemical synthesis.

EXAMPLE 1

Preparation of 7-t6-chloro-3-pyridinyl)-1-azabicyclo[2,2,1]heptane

20 Step A: <u>6-Chloro-N-methoxy-N-methyl-3-pyridine carboxamide</u>

To a suspension of 6-chloronicotinyl chloride (24.0 g, 0.136 mol) and N.O-dimethylhydroxylamine hydrochloride (14.0 g, 0.144 mol) in 200 mL of dichloromethane was added triethylamine (36.3 g, 0.358 mol) at 0°C. After complete addition, the reaction was warmed to room temperature and stirred for 2 h. Water was added and the mixture was extracted with dichloromethane. The organic layers were dried over MgSO₄, filtered and concentrated. Ether was added and the triethylamine hydrochloride salts were removed by filtration. Concentration of the ether solution gave 27.0 g of a yellow oil, sufficiently pure for the next step. ¹H NMR (CDCl₃) δ 8.78 (s,1H), 8.04 (d,1H), 7.40 (d,1H), 3.56 (s,3H), 3.40 (s,3H).

15

20

25

30

35

18

Step B: (6-Chloro-3-pyridinyl)(1-methyl-4-piperidinyl) methanone

Chloro(1-methyl-4-piperidinyl) magnesium (prepared from 4-chloro-1-methyl piperidine (17.75 g. 0.133 mol) in 300 mL of tetrahydrofuran and magnesium (3.4 g. 0.140 mol) according to the procedure in *J. Med. Chem.* (1965) 8, 829), was added to a solution of the product of Step A (13.3 g. 0.066 mol) in 50 mL of tetrahydrofuran at 0°C. After addition was complete, the reaction was warmed to room temperature. After 1 h, dilute HCi solution was added and the mixture was extracted with ethyl acetate. The combined extracts were dried over MgSO₄, filtered and concentrated. The crude material was triturated with hexane to give 9.8 g of an off-white solid, m.p. 104.5-106°C. ¹H NMR (CDCl₃) 8 8.90 (s.1H), 8.20 (d.1H), 7.45 (d.1H), 3.20-3.05 (m.1H), 2.95 (d.2H), 2.31 (s.3H), 2.20-2.00 (m.2H), 2.00-1.80 (m.4H).

Step C: (6-Chlore-3-pyridinyl)(4-piperidinyl) methanone hydrochloride

To a solution of the product of Step B (6.28 g, 0.026 mol) in 100 mL of 1.2-dichloroethane was added 1-chloroethyl chloroformate (9.0 g, 0.063 mol). The mixture was heated at reflux for 3-4 h. After cooling to room temperature, water was added and the mixture was extracted with dichloromethane. The combined extracts were dried over MgSO₄, filtered, and concentrated to give 8.0 g of a yellow solid. The crude solid was dissolved in methanol and heated at reflux for 30 min. After cooling to room temperature, the solvent was removed and the solids were triturated with ether to give 5.5 g (6-chloro-3-pyridinyl) (4-piperidinyl) methanone hydrochloride.

Step D: 2-Chloro-5-[chloro(4-piperidinyl)methyl] pyridine

Sodium borohydride (0.81 g, 0.021 mol) was added portionwise to a suspension of the product of Step C (5.5 g, 0.021 mol), in 50 mL of methanol. After complete addition, the reaction was stirred for 10 min and concentrated. The residue was suspended in 60 mL of dichloromethane/pyridine (5:1). Excess thionyl chloride (ca. 4-mL)-was-added. After-l-h, the-mixture-was-concentrated. Water-was-added-and-the-mixture was neutralized with potassium carbonate and extracted with ethyl acetate. The combined organic layers were dried over MgSO₄, filtered, and concentrated to give 2.6 g of an off-white solid.

Step E: 7-(6-Chloro-3-pyridinyl)-1-azabicyclo[2,2,1] heptane

Potassium *t*-butoxide (2.5 g, 0.022 mol) was added to a suspension of the product of Step D (2.6 g, 0.009 mol) in 50 mL of tetrahydrofuran. The mixture was heated at reflux for 48 h. After cooling to room temperature, water was added and the mixture was extracted with ethyl acetate. The combined extracts were dried over MgSO₄, filtered, and concentrated. Chromatography on silica gel (9:1 chloroform/methanol) afforded 1.3 g of a yellow solid, m.p. 77-78°C. ¹H NMR (CDCl₃) δ 8.42 (s.1H), 7.75 (d.1H), 7.27 (s.1H), 3.75 (s.1H), 3.05 (dt.1H); 2.90 (d.1H), 2.70-2.60 (m.2H), 2.50-2.40 (m.1H), 1.9-1.7 (m.1H), 1.45-1.30 (m.2H), 1.20-1.10 (m.1H).

15

20

30

19

EXAMPLE 2

Prenaration of 7-(3-pyridinyl)-1-azabicyclo[2,2,1]heptane

To a solution of 7-(6-chloro-3-pyridinyl)-1-azabicyclo [2.2.1]heptane (100.0 mg, 0.43 mmol) in methanol was added a catalytic amount of 5% palladium on carbon and a balloon containing hydrogen gas. After stirring overnight, the mixture was filtered through a pad of Celite[®] and concentrated. The residue was taken up in dichloromethane, washed with saturated sodium bicarbonate solution, dried over MgSO₄, filtered, and concentrated to give 55 mg of a brown oil. ¹H NMR (CDCl₃) 8 8.65 (s.1H), 8.50 (d.1H), 7.80 (d.1H), 7.25 (d.1H), 3.81 (s.1H), 3.10-3.00 (m.1H), 3.00-2.95 (m.1H), 2.80-2.60 (m.2H), 2.50-2.40 (m.1H), 1.90-1.80 (m.1H), 1.45-1.10 (m.2H).

EXAMPLE 3

Preparation of 7-(5.6-dichloro-3-pyridinyl)-1-azubicyclo[2,2,1]heptane Step A: 5,6-Dichloro-N-methoxy-N-methyl-3-pyridine carboxamide

To a solution of 5.6-dichloronicotinic acid (10.0 g, 0.052 mmol) and triethylamine (15.2 mL, 0.109 mmol) in dichloromethane was added oxalyl chloride (6.94 g, 0.05 mmol). After the gas evolution ceased, N.O-dimethylhydroxylamine hydrochloride (5.34 g, 0.055 mmol) was added and the mixture was stirred overnight. Saturated sodium bicarbonate solution was added and the mixture was extracted with dichloromethane. The combined organic layers were dried over MgSO₄, filtered, and concentrated to give 9.1 g of a brown solid. ¹H NMR (CDCl₃) δ 8.68 (s.1H), 8.16 (s.1H), 3.59 (s.3H), 3.40 (s.3H)

Step B: 7-(5.6-Dichloro-3-pyridinyl)-1-azabicyclof2.2.11heptane

The product of Step A was manipulated according to Steps B-E of Example 1 to give-0.95-g-of-a-waxy-solid, m.p. 30-32²C. ¹H NMR (CDCl₃) δ.8.31 (s.1H), 7.88 (s.1H), 3.73 (s.1H), 3.05 (t.1H), 2.95-2.90 (m.1H), 2.7-2.6 (m.2H), 2.5-2.4 (m.1H), 1.9-1.8 (m.1H), 1.45-1.30 (m.2H), 1.3-1.2 (m.1H).

By the procedures described herein, the following compounds of Tables 1 to 23 can be prepared. The compounds in Table 1, line 1 can be referred to as 1-1, 1-2, 1-3, 1-4 and 1-5 (as designated by line and column). All the other specific compounds covered in these Tables can be designated in an analogous fashion. The following abbreviations have been used in Tables 1-23: Me = methyl, Et = ethyl, iPr = isopropyl, nPr = n-propyl and Bn = benzyl.

20 <u>Table 1</u>

$$\begin{array}{c|c}
(CH_2)_n & \xrightarrow{1} & \xrightarrow{R^2} \\
1 & & & \\
1 & & & \\
1 & & & \\
2 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & & & \\
1 & &$$

		_					
		1			OLUMN		
•			1	2	3	4	5
1	$n=0; q=2; R^2=H; R^3=H;$	R1=	F	CI	Br	СН3	н
<u> </u>	$n=1$; $q=2$; $R^2=H$; $R^3=H$;	R1=	F.	CI	. Br	СН3	Н.
3	$n=0$; $q=2$; $R^2=5$ -C1; $R^3=H$;	R1=	F	Cı	Br	СН3	н.
4	n=1; q=2; R ² =5-C1; R ³ =H;	R ¹ =	F	Cı	Br	CH ₃	н
5	$n=0$; $q=2$; $R^2=5$ -F; $R^3=H$;	RI=	F	CI	Br	CH ₃	н
6	$n=1$; $q=2$; $R^2=5-F$; $R^3=H$;	R!=.	F	Cl	Br	CH ₃	Н
7	$n=0$; $q=2$; $R^2=5$ -CN; $R^3=H$;	R1=	F	Cl	Br	CH ₃	н
8	$n=1$; $q=2$; $R^2=5$ -CN; $R^3=H$;	RI=	F	CI	Br	CH ₃	н
9	$n=1$; $q=1$; $R^2=H$; $R^3=H$;	R ¹ =	F	CI	Br	CH ₃	Н
10	$n=1$; $q=1$; $R^2=5$ -C1; $R^3=H$;	R1=	F	Cl	Br	СН3	Н
11	n=1; q=1; R ² =5-F; R ³ =H;	R ¹ ≃	F	CI	Br	CH ₃	н
12	$n=1$; $q=1$; $R^2=5$ -CN; $R^3=H$;	R ¹ =	F	CI	Br	CH ₃	н
13	$n=0$; $q=2$; $R^2=H$; $R^3=2$ - CH_3 ;	R1=	F	CI	Br	CH ₃	н
14	$n=1$; $q=2$; $R^2=H$; $R^3=2$ - CH_3 ;	R ¹ =	F	CI	Br	CH ₃	н
15	$n=1$; $q=1$; $R^2=H$; $R^3=2$ - CH_3 ;	R1=	F	CI	Br	CH ₃	н
16	n=0; q=2; $R^2=H$; $R^3=3-NO_2$;	R1=	F	CI	Br	CH ₃	н
17	$n=1$; $q=2$; $R^2=H$; $R^3=3$ - NO_2 ;	· R1=	F	CI	Вг	CH ₃	Н
18	$n=1$; $q=1$; $R^2=H$; $R^3=3-NO_2$;	R1=	F	CI	. Br	CH ₃	н
19	$n=0$; $q=2$; $R^2=H$; $R^3=3$ -CN;	R 1 =	F	CI	Br	CH ₃	H,
20	$n=1$; $q=2$; $R^2=H$; $R^3=3$ -CN;	R1=	F	CI	Br	CH ₃	Н
21	$n=1$; $q=1$; $R^2=H$; $R^3=3$ -CN;	R!=	F	Cı	Br	CH ₃	Н

21 <u>Table 2</u>

$$\begin{array}{c|c}
R^{\frac{3}{3}} & (CH_{2})_{0} & \frac{4}{3} & \frac{5}{6} & R^{1} \\
& & & & & & & \\
2 & & & & & & & \\
\end{array}$$

				С	OLUMN		
			ı İ	2		1	5
22	n=(); q=2; R ³ =Cl;	R1=	F	Ci	Br	CH3	н
23	$n=1$; $q=2$; $R^3=C1$;	R1=	F	CI	Br	CH ₃	н
24	$n=1$; $q=1$; $R^3=C1$;	. R1=	F	Cl	. Br	CH3	н
25	n=0; q=1; R3=Br;	R1=	F	CI	Br	CH ₃	н
26	$n=1$; $q=2$; $R^3=Br$;	R ^l =	F	CI '	Br	CH ₃	н
27	$n=1$; $q=1$; $R^3=Br$;	R ^I =	F	CI	Br	CH ₃	·H
28	n=0; q=2; R ³ =OH;	R ^I =	F	CI	Br	CH ₃	Н
29	n=1: q=2: R ³ =OH:	R ¹ =	F	CI	Br	CH ₃	Н
30	$n=1$; $q=1$; $R^3=OH$;	R!=	F	Cl	Br	CH ₃	H
31	$n=0; q=2; R^3=NH_2;$	R1=	F	CI	Br	CH ₃	Н
32	$n=1$: $q=2$: $R^3=NH_2$:	R1=	F	CI	Br	CH ₃	H
33	$n=1$; $q=1$; $R^3=NH_2$;	R!=	F	CI	Br	CH ₃	н
34	$n=0; q=2; R^3=CN;$	R1=	F	CI	Br	CH ₃	н
35	$n=1$; $q=2$; $R^3=CN$;	R1=	F	CI	Br	CH ₃	н
36	$n=1$; $q=1$; $R^3=CN$;	R1=	F	CI	Br	CH ₃	н
37	$n=0$: $q=2$: $R^3=CH_3$:	R ! =	F	CI	Br	CH3	н
38	$n=1$; $q=2$; $R^3=CH_3$;	R ! =	F	CI	Br	CH3	н
39	$n=1$; $q=1$; $R^3=CH_3$;	₁ =	F	CI	Br	CH ₃	Н
40	n=0; q=2; R3=CO ₂ Me;	R1 =	F	CI	Br	CH ₃	н
4;	$n=1$; $q=2$; $R^3=CO_2Me$.	R1=	F	CI	Br	CH ₃	н
42	$n=1$; $q=1$; $R^{3}=CO_{2}Me$.	R1=	F	CI	Br	CH3	Н
43	$n=0; q=2; R^3=CO_2H;$	R ¹ = '	F	CI	Br	CH ₃	Н
44	n=1; q=2; R3=CO ₂ H;	R1=	F	CI	Br	CH3	Н
45	$n=1; q=1; R^3=CO_2H$.	R ¹ =	F	: CI	Br	CH:	<u> </u>

Table 3

$$(CH_{2^{l_{n}}}) \xrightarrow{\frac{1}{2}} (CH_{2^{l_{q}}}) \xrightarrow{\frac{1}{2}} (CH_{2^{l_{q}}}) \xrightarrow{\frac{1}{2}} (CH_{2^{l_{q}}})$$

		-					
		Ĺ		CC	DLUMN		
		Į		2	3	4	5
46	n=0; q=2; R ³ =3·NO ₂ ;	R1=	F	CI	Br	CH ₃	н
47	n=1; q=2; R ³ =3·NO ₂ ;	R ¹ =	F	CI	Br	CH ₃	н
48	n=1; q=1; R ³ =3-NO ₃ ;	R1=	F	CI '	Br	CH ₃	Н
4 9	n=0; q=2; R ³ =3'-CN;	R1=	F	CI	Βr	СН3	Н
50	$n=1$; $q=2$; $R^3=3$ -CN;	R!=	F	CI	Br	CH ₃	Н
51	$n=1$; $q=1$; $R^3=3$ -CN;	R1=	F	CI	Br	CH ₃	Н
52	$n=0$; $q=2$; $R^3=3-NO_2$, $2-NH_2$;	R ¹ =	F	CI	Br	CH ₃	Н
53	$n=1$; $q=2$; $R^3=3-NO_2$, $2-NH_2$;	R1= .	F	CI	Br	CH ₃	Н
54 /**	$n=1$; $q=1$; $R^3=3-NO_2$, $2-NH_2$;	R1=	F	Cl	Br	CH ₃	н
55	$n=0$; $q=2$; $R^3=3-NO_2$, 2NHMe;	R ¹ =	F	CI	Br	СН3	Н
56	$n=1$; $q=2$; $R^3=3-NO_2$, $2-NHMe$;	R ! =	F	CI	₿r	CH ₃	н
57	n=1; q=1; R3=3-NO ₂ , 2-NHMe;	R ! =	F	, CI	Br	CH ₃	н
58	$n=0$; $q=2$; $R^3=3-NO_2$, $2-CH_3$;	R ^l =	F	Cl	Br	CH ₃	н
59	n=1; q=2; R ³ =3-NO ₂ , 2-CH ₃ ;	R1=	F	CI	Br	CH ₃	н
60	$n=1$; $q=1$; $R^3=3-NO_2$, 2^3-CH_3 ;	R 1 =	F	CI	Br	CH ₃	н
61	n=0; q=2; R ³ =3-CN, 2-CH ₃ ;	R1=	F	CI	Br	CH ₃	Н
62	$n=1$: $q=2$; $R^3=3$ -CN, 2-CH ₃ ;	R1=	F	CI	Br	CH ₃	н
63	$n=1$; $q=1$; $R^3=3$ -CN, 2-CH ₃ ;	R ! =	F	CI	Br	CH ₃	н

Table 4

$$- \underbrace{ (CH_2)_n}_{N} \underbrace{ R^1}_{R^4}$$

			COLUMN			
			1	2	3	1
<i>د</i> ٠	R4=H; n=1;	R ¹ =	CI	Br	СНЗ	н
65	$R^4=H; n=2;$	R ¹ =	Cl	Br	СН3	н
66	R ² =CH ₃ ; n=1;	R ! =	CI	Br	CH ₃	н
57	$R^4 = CH_3; n=2;$	R1=	Ci	Br	CH ₃	н
68	R4=Bn; n=1;	R ¹ =	Cl	Br	CH ₃	н
59	$R^2 = B n; n = 2;$	R1=	C!	Br	CH:	<u> ⊢н</u>

23

Table 5

	•
76	R4=H; n=1;
71	R4=H; n=2;
72	:R4=CHp: n=1;
73	R4=CH3: n=2:
7≐	R4=Bn; n=1;
7.5	R4=Bn; n=2;

١	COLUMN				
ĺ	1 2 3 4				
	CI	Br	CH ₃	н	
	CI	Br	CH ₃	H	
	CI	Br	CH ₃	Н	
	CI	Br	CH ₃	н	
	CI	Br	CH ₃	н	
	CI	Br	CH3	н	

Table 6

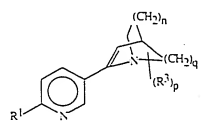
 $R^{1} =$ $R^{1} =$ $R^{1} =$

7 :	R ⁴ =H, n=1;
; -	R ⁴ =H; n=2;
78	R4=CH;: n=1:
7-4	8 ⁴ =CH ₃ , n=2;
5.1	R4=Bn: n=1;
S 1	84=85: 5=2:

	COLUMN					
	.,	2	3 -			
R1=	Cl	Br	CH;	н		
R 1 =	CI	Вг	CH ₃	Н		
R!=	Cl	Br	CH ₃	н		
R1=	C!	Br	CH3	H		
R1=	Ci	Br	CH3	н		
R1=	C:	8.	CH:	Н		

24

Table 7



82	n=1; q=1
\$3	n=1, q=2
c •	n=() ==3

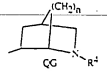
	COLUMN							
	ı	2	3	1	.5			
R1=	F	CI '	Br	CH3	Н			
R1=	F	Cl	Br	CH ₃	Н			
R!=	F	CI	Br	CH ₃	Н			

Key for Tables 8-23

where
$$Q^1 =$$

$$R^3 = \frac{1}{N} = (CH_2)_n$$

$$QA$$



25

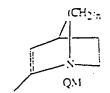


Table S

		ļ		C	OLUN	IN		
	$Q^{1} = QA$		1	2	3	4	5	
\$5	$Z=0$; $V=N$; $Y=CR^{T}$; $X=N$; $R^{T}=H$; $n=1$;	R1=	F	CI	Br	CH ₃	NH ₂	
86	Z=0: V=N: Y=CR ¹ : N=N: R ³ =H: n=2:	R1=	F	CI	Br	CH ₃	NH ₂	
.87	Z=O: V=N: Y=CR ¹ : N=N: R ³ =2 -CH ₃ : n=1:	R 1=	F	CI	Br	CH ₃	NH ₂	
SS	Z=0: V=N: Y=CR ¹ : X=N: R ³ =2-CH ₃ : n=2:	R1=	F	CI	Br	CH ₃	NH ₂	
89	Z=O: $V=N$: $Y=CR^{\frac{1}{2}}$: $X=N$: $R^{\frac{3}{2}}=4$ -OCH ₃ : $n=1$:	R!=	F	CI	Вг	CH ₃	NH ₂	ĺ
90	Z=0: V=N: Y=CR ¹ : X=N: R ³ =4-OCH ₃ : n=2:	R1=	F	CI	Вг	CH ₃	NH ₂	ĺ
91	Z=0; $V=N$; $Y=CR^{-1}$; $N=N$; $R^{-3}=4$ -Cl; $n=1$;	R1=	F	CI	Br	CH ₃	NH ₂	
92	Z=0: V=N: Y=CR ¹ : N=N: R ³ =4 ¹ -CI: n=2:	R1=	F	CI	Br	CH ₃	NH ₂	
93	Z=0: $V=N$: $Y=CR^{1}$: $X=N$: $R^{3}=4$ - CH_{3} : $n=1$:	R!=	F	CI	Br	CH ₃	NH2	
94	Z=0: V=N: Y=CR ¹ : X=N: R ³ =4-CH ₃ : n=2:	R1=	F	CI	Br	CH ₃	NH ₂	
95	Z=S; V=N; Y=CR ¹ ; X=N; R ³ =H; n=1;	R1=	F.	CI	Br	CH ₃	NH ₂	
96	Z=S: $V=N$: $Y=CR^{1}$: $X=N$: $R^{3}=H$: $n=2$:	R1=	F	CI	Br	CH ₃	NH ₂	
97	Z=S: N=N: Y=CR ¹ : N=N: R ³ =2-CH ₃ : n=1;	R1=	1:	CI	Br	CH ₃ .	NH ₂	
98	Z=S: $V=N$: $Y=CR^{-1}$: $N=N$: $R^{-3}=2$ -CH ₃ : $n=2$:	R1=	l:	CI	Br	CH ₃	NH ₂	
99	Z=S; V=N; Y=CR ¹ ; N=N; R ³ =4-OCH ₃ ; n=1;	R!=	F	CI	Br	CH ₃	NH ₂	
100	Z=S; V=N; Y=CR ¹ ; N=N; R ³ =4-0CH ₃ ; n=2;	R-1=	-F-	- -CI	Br	CH ₃	NH ₂	-
101	Z=S; V=N, Y=CR ¹ ; N=N; R ³ =4 ¹ ·Cl; n=1;	R1=	F	CI	Br	CH ₃	NH ₂	
102	Z=S: V=N: Y=CR ¹ : X=N: $R^3=4$ -CI: $n=2$:	R1=	Į.	CI	Br	CH ₃	NH ₂	
103	Z=S: V=N: Y=CR ¹ ; N=N: R ³ =4 -CH ₃ : n=1:	R1=	15	CI	Вг	CH ₃	NH ₂	
!(14	Z=S: N=N: Y=CR ¹ : N=N: R ³ =4 -CH ₃ : n=1:	R1=	F	CI	Br	CH ₃	NH ₂	
105	Z=N: V=N, Y=CR ¹ : N=O; R ³ =H; π=1;	R1=	F	CI	Br	CH ₃	NH ₂	
106	•	R!=	F	C	Br	CH ₃	NH ₂	
107	Z=N, V=N, Y=CR ¹ ; N=O; R ³ =2-CH ₃ ; n=1;	R!=	F	c	Br	СН3	NH ₂	
108	Z=N: V=N: Y=CR ¹ , N=O: R ² =2 ¹ -CH ₃ : n=2;	R1=	F	c	Br	CH ₃	NH ₂	
109	Z=N: V=N: Y=CR ¹ , X=O: R ² =4 ¹ -0CH ₃ : n=1;	R!=	F	С	i Br	CH ₃	NH2	
П	V ZHN, V=N, Y=CR ³ ; N=O; R ³ =4 ³ (OCH ₃ ; n=2)	R1=	F	C	Br	CH ₃	NH2	
1:1	$Z=N, V=N; Y=CR^{\frac{1}{2}}; X=O; R^{\frac{1}{2}}=4^{\frac{1}{2}}; C_{+}, n=1;$	R:=	F	С	i Br	CH;	NH ₂	
113	$Z = X_1 V = X_2 V = CR^{-1} (X = O(R)^{\frac{1}{2}} + 4^{\frac{1}{2}} + C(n = 2)$	R!=	i i	C	i Br	CH ₃	NH ₂	.
113	3 Z=N: V=N: Y=CR ¹ : N=O: R ² =4 (CH ₂ : n=1)	R!=	; į F	C	I Br	CH ₃	NH ₂	.

TO SHAME SALVENIES AND SHOW CONTRACTOR

Software the Market Warner

die die die

WO 95/03306

PCT/US94/08404

114	Z=N: V=N: Y=CR ¹ : X=O: R^3 =4-CH ₃ : n=2:	RI=	F	CI	Br	CH ₃	NH ₂	
115	Z=CH; V=N; Y=C(R^{1})=N; N=CH; R^{3} =H; n=1;	R¹=	F	CI	Br	СН3	н	
116	Z=CH; V=N; Y=C(R^{T})=N; X=CH; R^{3} =H; n=2;	R¹=	F	CI	Br	CH ₃	Н	
117	Z=CH: V=N: Y=C(R ¹)=N: X=CH: $R^3=2$ -CH ₃ : n=1:	R1= -	F	CI	Br	CH ₃	Н	
118	Z=CH: V=N: Y=C(R ¹)=N: X=CH: R^3 =2-CH ₃ : n=2:	R1=	F	CI	Br	CH ₃	Н	
119	Z=CH; V=N; Y=C(R ¹)=N; X=CH; R ³ =4-OCH ₃ ; n=1;	RI=	F	CI	Br	CH ₃	Н	
120	Z=CH; V=N; Y=C(R ¹)=N; X=CH; R ³ =4-OCH ₃ ; n=2;	R1=	F	CI	Br	СН3	Н	
121	$Z=CH; V=N; Y=C(R^{T})=N; X=CH; R^{3}=4'-Cl; n=1;$	R1=	F	CI	Вг	СН3	Н	
122	Z=CH; V=N; Y=C(R 1)=N; X=CH; R 3 =4-Cl; n=2;	R¹=	F	CI	Br	СН3	Н	
123	Z=CH; V=N; Y=C(R ¹ :=N; X=CH; R ³ =4'-CH ₃ ; n=1;	R1=	F	CI	Br	СН3	н	
124	Z=CH; V=N; Y=C(R ¹)=N; X=CH; R ³ =4-CH ₃ ; n=2;	R1= .	F	CI	Br.	CH ₃	H.	
125	Z=CH; V=N; Y=CR ¹ ; X=S; R ³ =H; n=1;	$R^1 =$	F	CI	Br	СН3	Н	
126	Z=CH: $V=N$: $Y=CR^{T}$: $N=S$: $R^{3}=H$: $n=2$:	R1=	F	CI	Br	CH ₃	н	
127	Z=CH; V=N; Y=CR 1 ; X=S; R 3 =2-CH $_{3}$; n=1;	K ¹=	F	CI	Br	CH ₃	н	
128	Z=CH: V=N: Y=CR ¹ : X=S: R^3 =2-CH ₃ : n=2;	R1=	F	CI	Br	СН3	н	
129	Z=CH: V=N; Y=CR ¹ ; X=S; R ³ =4-OCH ₃ ; n=1;	R1=	F	CI	Br	СН3	н	
130	Z=CH; V=N; Y=CR 1 ; X=S; R 3 =4-OCH $_{3}$; n=2;	R1=	F	Cı	Br	СН3	н	
131	Z=CH; V=N; Y=CR 1 ; X=S; R 3 =4 2 -Cl; n=1;	R1=	F	CI	Br	CH ₃	н	
132	Z=CH: $V=N$: $Y=CR^{-1}$: $X=S$: $R^{-3}=4$ -CI: $n=2$:	$R^{i}=$	F	CI	Br	CH ₃	н	
133	Z=CH: V=N: Y=CR ¹ : $X=S$: $R^3=4$ -CH ₃ : $n=1$:	R!=	F	CI	Br	СН3	Н	
134	Z=CH; $V=N$; $Y=CR^{-1}$; $N=S$; $R^{3}=4$ -CH ₃ ; $n=2$;	R ^I =	F	CI	Br	СН3	н	1
135	Z=0: V=N: Y=CR ¹ : X=CH: R ³ =H: n=1;	R1=	F	CI	Вг	CH ₃	Н	
136	Z=0: V=N; Y=CR ¹ : X=CH; R ³ =H; n=2;	R1=	F	CI	Br	CH ₃	н	Ì
137	Z=0: V=N: Y=CR ¹ : X=CH: R^3 =2-CH ₃ : n=1:	R1=	F	CI	Br	CH ₃	н	
138	Z=O: V=N: Y=CR ¹ : X=CH: R^3 =2-CH ₃ : n=2:	$R^1 =$	F	CI	Br	CH ₃	н	İ
139	Z=0: V=N: Y=CR ¹ : N=CH: R ³ =4-OCH ₃ : n=1:	R1=	F	CI	Br	CH ₃	Н	
140	Z=0: $V=N$: $Y=CR^{T}$: $N=CH$: $R^{3}=4$ -OCH ₃ : $n=2$:	R1=	F	CI	Br	СН3	Н	
141	Z=O; $V=N$; $Y=CR^{-1}$; $X=CH$; $R^{-3}=4^{-1}-CI$; $n=1$;	R ! =	F	CI	Br	CH ₃	н	j
142	$Z=0: V=N: Y=CR^{T}: X=CH: R^{T}=4^{T}-CI: n=2:$	R1=	F	CI	Br	CH ₃	н	Ì
143	$S = Z = O: V = N: V = CR^{\frac{1}{2}}: N = CH: R^{\frac{3}{2}} = 4^{\frac{1}{2}} - CH_3: n = 1;$	R1=	F	C	Br	CH ₃	H	i
144	$V = Z=0$; $V=N$; $Y=CR^{-1}$; $N=CH$; $R^{-3}=4$ - CH_3 ; $n=2$;	R 1 =	F	c	l Br	СН3	Н	į
143	$Z=N: V=CR^2: Y=C(R^3)=N: X=CH: R^2=H: R^3=H: n=1:$	R1=	F	c	ı Br	CH ₃	н	
140	$S = Z = N; V = CR^{2}; Y = C(R^{\frac{1}{2}}) = N; X = CH; R^{2} = H; R^{3} = H; n = 2;$	R1=	F	С	l Br	CH ₃	н	
141	$Z=N, V=CR^2, Y=C(R^4)=N; X=CH; R^2=H; R^3=2^4-CH_3; n=1;$	RI=	F	c	ı Br	CH ₃	Н	
143	$S = Z=N; V=CR^2; Y=C(R^3)=N; X=CH; R^2=H; R^3=2^3-CH_3; n=2;$	R!=	F	С	i Br	CH ₃	H	
14	 Z=N: V=CR²: Y=C(R³:=N: N=CH: R²=H: R³=4 -OCH₂: n=1: 	: R!=	F	c	ı Br	CH ₃	Н	
15	0 Z=N; V=CR ² , Y=C(R ³ ,=N; N=CH; R ² =H; R ³ =4 ³ -OCH ₃ ; n=2 ³	: R1=	: F	:	:	CH ₃	i	
15		R!=	i		!	· CH3	i	
			•	•	•		•	

BEST AVAILABLE COPY PCT/US94/08404

27

152	$Z=N$; $V=CR^{\frac{1}{2}}$; $Y=C(R^{\frac{1}{2}})=N$; $X=CH$; $R^{\frac{1}{2}}=H$; $R^{\frac{3}{2}}=4$; CI ; $n=2$;	R1=	F	CI	Br	сн3	н	
153	$Z=N$: $V=CR^2$: $Y=C(R^3)=N$: $X=CH$: $R^2=H$: $R^3=4$ - CH_3 : $n=1$:	R ¹ =	F	CI	Br	CH ₃	н	
154	$Z=N$; $V=CR^2$; $Y=C(R^1)=N$; $X=CH$; $R^2=H$; $R^3=4$ - CH_3 ; $n=2$;	R!=	F	CI	Br	CH ₃	Н	İ
155	$Z=N: V=CR^2: Y=C(R^{\frac{1}{2}})=N: X=CH: R^2=F: R^3=H: n=1;$	R1=	F	Cl	Br	CH ₃	н	
156	$Z=N: V=CR^2: Y=C(R^4:=N: X=CH: R^2=F: R^3=H: n=2:$	R1=	F	Cl	Br	CH ₃	н	
157	$Z=N: V=CR^2: Y=C(R^1)=N: X=CH: R^2=CI: R^3=H: n=1:$	R1=	F	Cl	Вг	CH ₃	н	
- 158	$Z=N$: $V=CR^2$: $Y=C(R^1)=N$: $X=CH$: $R^2=CH$: $R^3=H$: $n=2$:	R1=	F	CI	Br	CH ₃	н	
159	Z=CH: V=N: Y=NR ¹¹ : X=CH: R ³ =H: n=1:	R11=	CH ₃	Εt	iPr	nPr	н	
160	Z=CH: $V=N$: $Y=NR^{11}$: $X=CH$: $R^3=H$: $n=2$:	R11=	CH ₃	Ει	iPr	nPr	н	
161	$Z=CH; V=N; Y=NR^{11}; X=CH; R^3=2^{-}CH_3; n=1;$	R11=	CH ₃	Et	iPr	nPr	н	
162	Z=CH: $V=N$: $Y=NR^{11}$: $X=CH$: $R^3=2^{\frac{1}{2}}$ -CH3: $n=2$:	· R11=	CH ₃	Et	iPr .	nPr -	H '	
163	$Z=CH: N=N: Y=NR^{1,1}: X=CH: R^{3}=4 - OCH_3: n=1:$	$R^{11}=$	CH ₃	Εt	iPr	nPr	н	
164	Z=CH: V=N: Y=NR $^{1.1}$: X=CH: R 3 =4 4 -OCH3: n=2:	R11=	CH3	Ει	iPr	nPr	Н	
165	$Z=CH; N=N; Y=NR^{11}; X=CH; R^3=4-CI; n=1;$	R11=	CH ₃	Et	iPr	nPr	н	
166	Z=CH; V=N; Y=NR ¹¹ : X=CH; R^3 =4 -CI; n=2;	R11=	CH ₃	Eı	iPr	nPr	н	
167	Z=CH; V=N; Y=NR 11 ; X=CH; R 3 =4 -CH ₃ ; n=1;	R11=	CH ₃	Et	iPr	nPr	н	
168	Z=CH: V=N; Y=NR ¹¹ ; X=CH; $R^3=4$ -CH3; n=2;	R11=	CH	Et	iPr	nPr	Н	

Table 9

	_						
	,	COLUMN					
<u>Q1=Q3</u>		t	2	3	4	5	
169 Z=O; V=N; Y=CR ¹ ; X=N; R ³ =H; n=1;	R1=	Į:	Cl	Br	CH ₃	NH ₂	i
170 Z=O; V=N; Y=CR ¹ ; X=N, R ³ =H; n=2;	R1=	ħ	Cl	Br	CH3	NH ₂	
171 Z=O: $V=N$: $Y=CR^{-1}$: $X=N$: $R^{-3}=2$ -CH ₃ : $n=1$:	R1=	F	CI	Br	СН3	NH ₂	
172 Z=O; V=N; Y=CR ¹ ; X=N; R ³ =2 -CH ₃ ; n=2;	R1=	ŀ	CI	Br	CH ₃	NH ₂	-
173 Z=0; V=N; Y=CR ¹ ; N=N; R ³ =4-OCH ₃ ; n=1;	R1=	F	CI	Br	CH ₃	NH ₂	
174 Z=O; V=N; Y=CR ¹ ; N=N; R ³ =4 -OCH ₃ ; n=2;	R 1 =	F	CI	Br	CH;	NH ₂	
175 $Z=0$; $V=N$; $Y=CR^{1}$; $N=N$; $R^{3}=4$ -Ci; $n=1$;	R1=	F	CI	Br	CH:	NH2	
176 Z=0; V=N; Y=CR ¹ ; X=N; R ³ =4 -Ci; n=2;	R1=	F	CI	Br	CH ₃	NH ₂	!
177 Z=O, V=N; Y=CR ¹ ; X=N; R ³ =4-CH ₃ ; n=1;	R1=	F	CI	Br	CH ₃	NH ₂	
178 Z=0: V=N, Y=CR ¹ : N=N: R ² =4 -CH ₃ : n=2:	R1=	F	CI	Br	CH ₃	NH ₂	
179 Z=S, V=N; Y=CR ¹ ; X=N; R ³ =H; π=1;	R!=	F	CI	Br	CH ₃	NH ₂	
180 Z=S; V=N; Y=CR ¹ ; N=N; R ³ =H; π=2;	R1=	F	CI	Br	CH ₃	NH ₂	
181 Z=S; V=N, V=CR ¹ ; X=N; R ³ =2 ³ -CH ₃ ; n=1;	R!=	F	CI	Br	1	NH ₂	•
182 Z=S, V=N, Y=CR ¹ ; N=N; R ² =2-CH ₃ ; n=2.	R1=	; r	Ci	: Br	i	NH ₂	٠,
183 Z=S, V=N, Y=CR ¹ ; N=N, R ³ =4 ¹ -OCH ₃ ; n=1;	R1=	ŀ	CI	: ! Br	1	NH ₂	٠.
184 Z=S, V=N, Y=CR ¹ , N=N, R ² =4 -OCH ₃ ; n=2;	R!=	. F	C:	: S:		NH	٠,

185	Z=S: $V=N$: $Y=CR^{-1}$: $X=N$: $R^{-3}=4$ -CI: $n=1$:	R!=	F	CI	Br	CH ₃	NH2	
186	Z=S: $V=N$: $Y=CR^{-1}$: $X=N$: $R^{-3}=4$ -CI: $n=2$:	R1=	F.	CI	Br	CH ₃	NH	
187	Z=S: $V=N$: $Y=CR^{-1}$: $X=N$: $R^{-3}=4$ - CH_3 : $n=1$:	R ¹ =	F	Cı	Br	CH ₃	NH	
133	Z=S: $V=N$: $Y=CR^{-1}$: $X=N$: $R^{-3}=4$ - CH_3 : $n=1$:	R ! =	F	CI	Br	CH ₃	NH ₂	
189	$Z=N; V=N; Y=CR^{-1}; X=0; R^{-2}=H; n=1;$	R1=	F [.]	Cı	Br	CH ₃	NHa	
190	Z=N: $V=N$: $Y=CR^{-1}$: $X=0$: $R^{-2}=H$: $n=2$:	R ¹ =	F	CI	Br	CH ₃	NHa	
191	Z=N: $V=N$: $Y=CR^{1}$: $X=0$: $R^{3}=2$ -CH ₃ : $n=1$:	R!=	F	Cı	Br	CH ₃	NH ₂	
192	Z=N; V=N; Y=CR 1 ; X=0; R 3 =2 1 -CH $_{3}$; n=2;	R ¹ =	F	Cı	Br	CH ₃	NH ₂	
193	$Z=N; V=N; Y=CR^{T}; N=0; R^{S}=4-OCH_{3}; n=1;$	R1=	F	CI	Br	CH ₃	NH ₂	
194	Z=N; V=N; Y=CR ¹ ; X=O; \mathbb{R}^3 =4-OCH ₃ ; n=2;	R1=	F	CI	Br	CH ₃	NH ₂	
195	$Z=N; V=N; Y=CR^{-1}; X=0; R^{-3}=4 \cdot Cl; n=1;$	R1=	F	CI	Br	CH ₃	NH ₂	
196	Z=N: V=N: Y=CR ¹ : X=O: R^3 =4 -C1: n=2:	R !=	F.	CI.	Br	СН3	NH ₂	
197	Z=N; V=N; Y=CR ¹ ; X=O; $R^3=4$ -CH ₃ ; n=1;	R 1 =	F	CI	Br	СН3	NH ₂	
198	Z=N; V=N; Y=CR 1 ; X=O; R 3 =4 -CH $_{3}$; n=2;	R1=	F	CI	Br	CH ₃	NH ₂	
199	Z=CH; $V=N$; $Y=C(R^{1})=N$; $X=CH$; $R^{3}=H$; $n=1$;	R ^l =	F	CI	Br	CH ₃	н	
200	Z=CH; $V=N$; $Y=C(R^{1})=N$; $X=CH$; $R^{3}=H$; $n=2$;	R1=	F	CI	Br	CH ₃	Н	
201	Z=CH; V=N; Y=C(R ¹)=N; X=CH; R ³ =2-CH ₃ ; n=1;	R ¹ =	F	CI	Вг	СН3	н	
202	Z=CH; V=N; Y=C(R ¹)=N; X=CH; R ³ =2-CH ₃ ; n=2;	R1=	F	CI	Br	СН3	н	
203	Z=CH: $V=N$: $Y=C(R^{\frac{1}{2}})=N$: $X=CH$: $R^{\frac{3}{2}}=4$ -OCH ₃ ; $n=1$;	R1=	F	CI	Br	СН3	н	
204	Z=CH: V=N: Y=C(R ¹)=N: N=CH: R ³ =4 1 -OCH ₃ : n=2:	R1=	F	CI .	Br	СН3	н	
205	Z=CH; $V=N$; $Y=C(R^{1})=N$; $N=CH$; $R^{3}=4$ -Cl; $n=1$;	R1=	F	CI	Br	CH ₃	н	
206	Z=CH: $V=N$: $Y=C(R^{-1})=N$: $X=CH$: $R^{-2}=4$ -CI: $n=2$:	R 1 =	F	CI	Br	СН3	Н	
207	Z=CH; V=N; Y=C(R ¹)=N; X=CH; R ³ =4-CH ₃ ; n=1;	R ¹ =	F	CI	Br	CH ₃	н	
208	Z=CH; V=N; Y=C(R^{1})=N; X=CH; R^{3} =4-CH ₃ ; n=2;	R1=	F	CI	Br	CH ₃	н	
209	Z=CH;_V=N;_Y=CR ¹ ;_X=5;_R ³ =H;_n=1;	R ¹ =	F	- -CI-	Br—		Н	-
210	Z=CH: $V=N$: $Y=CR^{-1}$: $X=5$: $R^{-3}=H$: $n=2$:	R 1 =	F	CI	Br	СН3	н	
211	Z=CH: $V=N$: $Y=CR^{-1}$: $X=S$: $R^{-3}=2^{-1}$ -CH ₃ : $n=1$:	R 1 =	F	CI	Br	CH ₃	н	
212	Z=CH; V=N; Y=CR ¹ ; X=S; R ³ =2 1 -CH ₃ ; n=2;	R ¹ =	F	CI	Br	CH ₃	1	
213	Z=CH: $V=N$: $Y=CR^{-1}$: $X=S$: $R^{-3}=4$ -OCH ₃ : $n=1$:	R ! =	F	CI	Br	CH ₃	н	1
214	Z=CH: V=N: Y=CR ¹ : X=3: R ³ =4 -OCH ₃ : n=2:	R!=	F	CI	Br	CH:	н	
215	$Z=CH; V=N; Y=CR^{\frac{1}{2}}; X=S; R^{\frac{3}{2}}=4 \cdot Cl; n=1;$	R1=	F	CI	Br	CH:	н	į
216	Z=CH: V=N: Y=CR ¹ : X=S: R ³ =4 -Cl: n=2:	R ¹ =	F	CI	Br	СН	Н	
217	Z=CH: V=N: Y=CR ¹ : X=3. R ³ =4 ¹ -CH ₃ : n=1;	R1=	F	cı	Br	CH:	; Н	!
213	S Z=CH: V=N: Y=CR ¹ : X=S: R ³ =4 ² -CH ₃ : n=2:	R1=	F	C!	Br	СН	Н	i
211) Z=O: N=N, Y=CR ¹ : N=CH, R ² =H: n=1:	R1=	F	Cı	Br	•	Н	
229) Z=0: V=N: Y=CR ¹ : X=CH: R ³ =H: n=2:	R ! =	: F	C:	Br	CH	i	İ
22	I Z=0: V=N, Y=CR ¹ : N=CH: R ³ =2-CH ₃ : n=1:	R ⁱ =	F	C!	. Br	СН	1	!
22	2 Z=0: V=N: Y=CR ⁴ : N=CH: R ³ =2-CH ₃ : n=2:	R1=	: F	CI	Br	СН	3 H	
							-	•

						٠.	,	
223	$Z=O: V=N: Y=CR^{1}: X=CH: R^{3}=4 -OCH_{3}: n=1:$	R1=	F	CI	Br	CH ₃	н	
224	Z=O: V=N: Y=CR ¹ : X=CH: R^3 =4'-OCH ₃ : n=2:	R1=	F'	CI	Br	CH ₃	н	
225	Z=O: $V=N$: $Y=CR^{1}$: $X=CH$: $R^{3}=4$ -CI: $n=1$:	R1=	F	CI	Br ·	CH ₃	н	
226	Z=0; $V=N$; $Y=CR^{1}$; $X=CH$; $R^{3}=4$ -Cl; $n=2$;	R1=	F	CI	Br	CH ₃	н	
227	Z=0; V=N; Y=CR ¹ ; X=CH; R ³ =4-CH ₃ ; n=1;	R1=	F	Cl	Br	CH ₃	н	
228	Z=0; V=N; Y=CR ¹ ; X=CH; $R^3=4$ -CH ₃ ; n=2;	R1=	F	CI	Br	CH ₃	н	
229	$Z=N$; $V=CR^2$; $Y=C(R^4)=N$; $X=CH$; $R^2=H$; $R^3=H$; $n=1$;	R ¹ =	F	CI	Вr	CH ₃	н	
230	$Z=N$; $V=CR^2$; $Y=C(R^1)=N$; $X=CH$; $R^2=H$; $R^3=H$; $n=2$;	R1=	F	Cı	Br	СН3	Н	
231	$Z=N$: $V=CR^2$: $Y=C(R^1)=N$: $X=CH$: $R^2=H$: $R^3=2$ - CH_3 : $n=1$:	R1=	F	Ci	Br	CH ₃	Н	
232	$Z=N$; $V=CR^2$; $Y=C(R^1)=N$; $X=CH$; $R^2=H$; $R^3=2$ - CH_3 ; $n=2$;	R1=	F	CI	Br	CH ₃	Н	
233	$Z=N_1 V=CR^2$; $Y=C(R^1)=N_1 X=CH$; $R^2=H$; $R^3=4$ -OCH ₃ ; $n=1$;		F	CI	Br	CH ₃	Н	
234	$Z=N$; $V=CR^2$; $Y=C(R^1)=N$; $X=CH$; $R^2=H$; $R^3=4$ -OCH ₃ ; $n=2$;		F	CI	Br	CH ₃	Н	
235	$Z=N$; $V=CR^2$; $Y=C(R^1)=N$; $X=CH$; $R^2=H$; $R^3=4$ -CI; $n=1$;	R1=	F	CI	Br	CH ₃	н	
236	$Z=N$; $V=CR^2$; $Y=C(R^1)=N$; $X=CH$; $R^2=H$; $R^3=4$ -CI; $n=2$;	R1=	F	CI	Br	CH ₃	н	
237	$Z=N; V=CR^2; Y=C(R^1)=N; X=CH; R^2=H; R^3=4-CH_3; n=1;$	R!=	F	CI	Br	CH ₃	н	
238	$Z=N$; $V=CR^2$; $Y=C(R^1)=N$; $X=CH$; $R^2=H$; $R^3=4$ - CH_3 ; $n=2$;	R 1 =	F	CI	Br	CH ₃	Н	
239		Ŕ ¹ =	F	CI	Br	CH ₃	н	
240	$Z=N$; $V=CR^2$; $Y=C(R^1)=N$; $N=CH$; $R^2=F$; $R^3=H$; $n=2$;	R1=	F	CI	Вг	CH ₃	н	
241		R1=	F	CI	Br	CH ₃	н	
242		R 1 =	F	CI	Br	CH ₃	н	
243		R11	-	Ει	iPr	nPr	Η.	
244	$Z=CH; V=N; Y=NR^{11}; X=CH; R^3=H; n=2;$	RII	= CH	Et Et	iPr	nPr	Н	
245		RH	= CH	Et	iPr	nPr	Н	
246	$S = Z = CH; V = N; Y = NR^{11}; X = CH; R^3 = 2 - CH_3; n = 2;$	RII	= CH	3 Et	iPr	nPr	Н	
24	$Z = Z = CH_1 \cdot X = N_1 \cdot Y = NR \cdot \frac{11}{11} \cdot X = CH_1 \cdot R^{\frac{3}{2}} = 4 \cdot OCH_3 \cdot n = 1$:	R11	<u>= CH</u>	Et Et	iPr	nPr	<u> H</u>	_ -
24	$S = Z = CH; V = N; V = NR^{11}; N = CH; R^3 = 4 - OCH_3; n = 2;$	RII	= CH	3 Et	iPr	ηPr	H	
2.1	9 Z=CH: N=N: Y=NR ¹¹ : X=CH: R ³ =4-Cl: n=1;		= СН	3 Et	iPr	nPr	H	İ
25		R ^T		• 1	iPr	nPr	н	-
25	1 Z=CH: V=N: Y=NR ¹¹ : X=CH: R ³ =4 ² -CH ₃ : n=1:	R!	i= CH	3 Et	iPr	nPr	Н	ļ
25	2 Z=CH; V=N; Y=NR ¹¹ ; X=CH; R ² =4-CH ₃ ; n=2;	RI	I= CH	Et Et	i.Pr	nPr	Н	

Table 10

			COLUMN				
	<u> </u>	<u> </u>	1	2	3	4	5
253	Z=S, N=CH; N=CR $^{\frac{1}{4}}$; N=N, n=1;	R1= :	CH;	OCH ₃	NH2	Ci	Н
254	Z=S: N=CH: Y=CR ¹ : N=N, n=2.	R!=	CH;	OCH;	NH ₂	CI	н
255	Z=O: V=CH: Y=CR ¹ , X=N, n=1,	₽1=	CH ₃	OCH ₃	NH ₂	Ci	н :
256	Z=O, V=CH; Y=CR ³ , X=N, n=2;	R = ;	CH ₃	OCH ₃	ZH2	Ci	н

WO 95/03306

BEST AVAILABLE COPY PCT/US94/08404

30

	•				_		_	
257	Z=CH: V=S: Y=CR ¹ , X=N, n=1;	R ¹ =	СН3	осн ₃	NH ₂	CI	н	
258	Z=CH: V=S: Y=CR ¹ , X=N, n=2:	R1=	сн3	осн3	NH2	CI	н	
259	Z=CH: $V=0$: $Y=CR^{-1}$: $X=N$: $n=1$:	R ¹ =	сн3	OCH ₃	ŇH ₂	CI	н	
260	Z=CH: $V=0$: $Y=CR^{-1}$, $X=N$. $n=2$:	R ¹ =	сн3	OCH3	NH ₂	CI	н	
261	Z=CR ¹ ; V=CH; Y=O, X=N, n=1;	R1=	CH ₃	OCH ₃	NH2	CI	н	
262	Z=CR ¹ ; V=CH; Y=O, X=N, n=2;	$R^{1}=$	СН3	och ₃	NH2	Cl	н	
263	Z=CH; V=CH=CH; Y=CR ¹ ; X=N; n=1;	R!=	СН3	осн3	NH ₂	CI	н	í
264	Z=CH: V=CH=CH: Y=CR ¹ : X=N: n=2;	R1=	СН3	осн ₃	NH ₂	Cl	н	
265	Z=CH: \hat{V} =N: Y=CH=C(R ¹): X=N: n=1:	R1=	CH ₃	OCH ₃	NH ₂	Cl	н.	l
266	$Z=CH$; $V=N$; $Y=CH=C(R^{T})$; $X=N$; $n=2$;	R 1 =	CH3	OCH ₃	NH ₂	CI	Н	
267	$Z=CH$; $V=N$; $Y=C(R^T)=N$; $N=CH$; $n=1$;	R1= .	F	CI ·	Br .	CH3.	н	
268	$Z=CH_1V=N; Y=C(R^1)=N; X=CH; n=2;$	R1=	F	CI	Br	CH ₃	Н	
269	Z=CH: V=N; Y=CR ¹ ; X=S; π=1;	R1=	F	CI	Br	CH ₃	H·	
270	Z=CH; V=N; Y=CŘ ¹ ; X=S; π=2;	R!=	F	CI	Br	CH ₃	н	
271	Z=0; V=N; Y=CR ¹ ; X=CH; n=1;	R1=	F	Cı	Br	CH ₃	Н	
272	Z=O; V=N; Y=CR ¹ ; X=CH; π=2;	R1=	F	CI	Br	CH ₃	H .	
273	$Z=N; V=CH; Y=C(R^{1})=N; X=CH; n=1;$	R1=	F	CI	Br	СН3	Н	
27-1	$Z=N; V=CH; Y=C(R^{-1})=N; X=CH; n=2;$	R1=	F	CI	Br	CH ₃	Н	
275	$Z=CH; V=N; Y=C(R^{T})=CH; X=CH; n=1;$	R1=	F	CI	Br	CH ₃	н	
276	$Z=CH: V=N: Y=C(R^{T})=CH: X=CH: n=2;$	R1=	F	CI	Br	CH ₃	Н	
277	Z=CH; V=CH; Y=NR ¹¹ ; X=N; n=1;	RII=	CH ₃	C ₂ H ₅	Н	iPr	nPr	
278	Z=CH; V=CH; Y=NR ¹¹ ; X=N; n=2;	R11=	-	C ₂ H ₅	Н	iPr	aPr	
279	Z=CH; V=N; Y=NR ¹¹ ; X=CH; n=1;	R11=	-	C ₂ H ₅	н	iPr	nPr	ĺ
280	Z=CH; V=N; Y=NR ¹¹ ; N=CH; n=2;	RII=	CH-	C ₂ H ₅	Н	iPr	nPr	

Table 11

			COLUMN				
	<u>Q </u>		1	2	3	4	5
281	$Z=0$: $V=N$: $Y=CR^{T}$: $N=N$: $R^{T}=H$: $n=1$:	R1=	F	CI	8r	CH ₃	NH ₂
282	Z=0: $V=N$: $Y=CR^{T}$: $X=N$: $R^{4}=CH_{3}$: $n=1$:	R1=,	F	Cl	Br	сн3	NH ₂
283	1	R ! =	F	CI	Вг	CH ₃	NH ₂
284	Z=0; V=N; Y=CR ¹ ; X=N; R ² =H; n=2;	R1=	F	CI	Br	СН3	NH ₂
285	Z=0: $V=N$: $Y=CR^{T}$: $X=N$: $R^{T}=CH_{3}$: $n=2$:	R1=	F	CI	Br	CH ₃	NH ₂
286	Z=0; V=N; Y=CR ¹ ; X=N; R ² =Bn; n=2;	R1=	F	CI	Br	CH ₃	NH2
287	$Z=S: V=N: V=CR^{T}: X=N: R^{T}=H: n=1:$	R1=	F	CI	Br	CH ₃	NH ₂
- 288	Z=S: $V=N$: $Y=CR^{T}$: $N=N$: $R^{T}=CH_{3}$: $n=1$:	R!=	F	CI	Br	CH ₃	NH ₂

WO 95/03306

	•							
289	$Z=S$, $V=N$; $Y=CR^{\frac{1}{2}}$; $N=N$; $R^{\frac{1}{2}}=Sn$; $n=1$;	R1=	F	CI	Br	CH ₃	NH ₂	
290	$Z=S: V=N: Y=CR^{T}: N=N: R^{T}=H: n=2:$	R1=	F	CI	Br	CH_3	NH ₂	
291	$Z=S: V=N: Y=CR^{\frac{1}{2}}: X=N: R^{\frac{1}{2}}=CH_3: n=2:$	R ! =	F	CI	Br	CH_3	NH ₂	
292	Z=S: $V=N$: $Y=CR^{\frac{1}{2}}$: $N=N$: $R^{\frac{2}{3}}=Bn$: $n=2$:	R =	F	CI	Br	CH_3	NH ₂	
293	$Z=N: V=N: Y=CR^{1}: X=0: R^{4}=H: n=1:$	R 1 =	F	CI	Br	CH_3	NH ₂	
<u> 2</u> 94	$Z=N: V=N: Y=CR^{\frac{1}{2}}: X=0. R^{\frac{2}{3}}=CH_3: n=1:$	R ! =	F	CI	Br_	CH_3	NH ₂	
295	$Z=N : V=N: Y=CR^{\frac{1}{2}}: X=0: R^{\frac{1}{4}}=Bn: n=1:$	R!=	F	Cl	Br	CH_3	NH ₂	
296	$Z=N: V=N: Y=CR^{\frac{1}{2}}: N=0: R^{\frac{1}{4}}=H: n=2:$	R 1 =	F	CI	Br	CH ₃	NH ₂	
297	$Z=N: V=N: Y=CR^{\frac{1}{2}}, N=0: R^{\frac{1}{2}}=CH_{3}; n=2:$	R ^I =	F	Cl	Br	CH ₃	NH ₂	
298	$Z=N: V=N: Y=CR^{\frac{1}{2}}: N=0: R^{\frac{1}{4}}=Bn: n=2:$	R1=	F	CI	Br	CH ₃	NH ₂	
299	$Z=CH; V=N; Y=C(R^{\frac{1}{2}})=N; N=CH; R^{\frac{1}{2}}=H; n=1;$	R!=	CI	Br	CH3			
300	$Z=CH; V=N; Y=C(R^{T})=N, X=CH; R^{4}=H; n=2;$	R1=	CI	Br	CH ₃	Ì	İ	
301	$Z=CH; V=N; Y=C(R^{1})=N; X=CH; R^{4}=CH_{3}; n=1;$	R 1 =	CI	Br	CH ₃			•
302	Z=CH: $V=N$: $Y=C:R^{1}:=N$: $N=CH$: $R^{4}=CH_{3}$: $n=2$:	R1=	CI	Br	CH ₃			
303	Z=CH; V=N; Y=CR ¹ ; X=S; R ² =H; n=1;	R1 =	CI	Br	CH ₃			
304	$Z=CH; V=N; Y=CR^{\frac{1}{2}}; N=S; R^{\frac{2}{3}}=H; n=2;$	R1=	CI	Br	CH ₃			1.
305	·	R1=	CI	Br	CH ₃			-
306	-	R ^I =	CI	Br	CH ₃			1
307		R ! =	CI	Br	CH ₃		}	
308	Z=0: $V=N$; $Y=CR^{\frac{1}{2}}$; $X=CH$; $R^{\frac{4}{2}}=H$; $n=2$;	R1=	CI	Br	CH ₃			
309	•	R 1 =	CI	Br	CH ₃			-
310	•	R1=	CI	Br	CH ₃			1
311		R 1 =	CI	Br	CH ₃			1
313		R ! =	CI	Br	CH ₃			
313		R¹=	CI	Br	CH ₃			
31-	•	R1=	CI	Br	CH ₃			
31:		R ¹ =	CI	Br	CH ₃			
31)		R1=		Br	CH ₃			
31	•	R!=	!	Br	CH ₃			
34	$S = Z = CH, V = N; Y = CR^{\frac{1}{2}}; X = S; R^{\frac{2}{3}} = CH_{\frac{1}{3}}; n = 2;$	R1=	CI	Br	CH3		j	
31		R 1 =	CI	Br	CH3			
32	0 Z=0; N=N; Y=CR ³ , N=CH; R ² =H; n=2;	R ¹ =	CI	Br	CH:	;		
32	$(i - Z=0) N=N; N=CR^{\frac{1}{2}}; N=CH; R^{\frac{1}{2}}=CH_3; n=1;$	R ¹ =	CI	ļ Br	CH:	:		
3.	2 Z=0, N=N; Y=CR ³ , N=CH; R ² =CH ₃ ; n=2;	R ⁱ =	CI	Br	СН	;	Ì	
33	$3 = Z = N; V = CH; Y = C; R^{T} := N; N = CH; R^{T} = H; n = 1;$	K¹=	Ci	13:	СН	;		
33	24 Z=N; V=CH; Y=C,R ³ ;=N, N=CH, R ³ =H; n=2;	K¹=	: C!	Bi	СН	3		
33	25 Z=N, V=CH; Y=C(R ¹)=N, N=CH; R ² =CH ₂ ; n=1.	R1=	: C1	. B:	- CH	3	ĺ	
33	26 Z=N: V=CH: Y=C:R ⁴ :=N: N=CH: R ⁴ =CH ₂ : n=2:	81=	- C!	8	: CH	3		

PCT/US94/08404

32

327	Z=CH: V=N:	Y=NR11:	X=CH:	R ⁴ =H. n=1;
-----	------------	---------	-------	-------------------------

328 Z=CH: V=N: Y=NR¹¹: N=CH: R⁴=H, n=2;

329 Z=CH; V=N; Y=NR¹¹; X=CH; R⁴=CH₃, n=1;

330 Z=CH; V=N; Y=NR¹¹; X=CH; R⁴=CH₃, n=2;

RII=	СН3	iPr	Et	1
R11=	СН3	iPr	Et	
R11=	CH ₃	iPr	Εt	İ
RII=	CH ₃	iPr	Et	

Table 12

	$Q^1 = QE \qquad R^4 = CH_3$
331	$Z=CH; V=N; Y=C(R^{1})=N; X=CH; n=1;$
332	$Z=CH; V=N; Y=C(R^{T})=N; X=CH; n=2;$
333	Z=CH; V=N; Y=CR ¹ ; X=S; n=1;
334	Z=CH; V=N; Y=CR ¹ ; N=S; n=2;
335	Z=0; V=N; Y=CR ¹ ; X=CH; n=1;
336	Z=O; V=N; Y=CR ¹ ; X=CH; n=2;
337	$Z=N; V=CH; Y=C(R^{1})=N; X=CH; n=1;$
338	$Z=N; V=CH; Y=C(R^{T})=N; X=CH; n=2;$
339	Z=O; V=N; Y=CR ¹ ; X=N; n=1;
340	Z=O; V=N; Y=CR ¹ ; X=N; n=2;
341	$Z=S: V=N: Y=CR^{1}: X=N: n=1:$
342	$Z=S: V=N: Y=CR^{\frac{1}{2}}: X=N: n=2;$
343	$Z=CH$; $V=N$; $Y=NCH_3$; $X=CH$; $n=1$;
344	$Z=CH; V=N; Y=NCH_3; X=CH; n=2;$

_						
Į	COLUMN					
	1	2	3			
RI=	CI	Br	СН3			
RI=	CI	Br	СН3			
R1=	CI	Br	CH ₃			
R1=	Cl	Br	CH ₃			
R1=	CI	Br	CH ₃			
R1=	CI	Br	CH ₃			
R1=	CI	Br	CH ₃			
R1=	CI	Br	CH ₃			
R!=	CI	NH ₂	СН3			
$R^{1}=$	CI	NH ₂	СН3			
R1=	Cı	NH ₂	CH ₃			
R1=	CI	NH ₂	CH ₃			
R!=	CI	NH ₂	СН3			
$R^1 =$	CI	NH ₂	CH ₃			

Table 13

	Į	COLUMN				
$Q^{1}=QF_{1}$ $R^{4}=CH_{3}$		1	. 2	3	1	5
345 Z=S: V=CH: Y=CR ¹ : X=N, n=1:	R ! =	CH ₃	оснз	NH ₂	CI	н
346 Z=5; V=CH; Y=GR ⁴ ; X=N, n=2;	R1=	CH ₃	OCH;	NH2	CI	н
347 Z=O: V=CH: Y=CR ¹ , X=N, n=1:	R1=	CH ₃	OCH ₃	NH ₂	CI	н
348 Z=O: V=CH: Y=CR ¹ , X=N, n=2:	R1=	CH ₃	OCH ₃	NH ₂	CI	н
349 Z=CH; V=S; Y=CR ¹ , N=N, n=1;	R ! =	CH ₃	OCH ₃	NH ₂	CI	н
350 Z=CH; V=S; Y=CR ¹ , X=N, n=2;	R 1 =	CH ₃	осн _з	NH ₂	CI	Н
351 Z=CH; N=O; Y=CR ¹ , X=N, n=1;	R ! =	CH ₃	OCH ₃	NH ₂	CI	Н
352 Z=CH; N=O; Y=CR ¹ , N=N, n=2;	R1=	CH ₃	OCH;	NH2	CI	н
353 Z=CR ¹ , V=CH; Y=O, X=N, n=1;	R1=	CH ₃	OCH ₃	NH ₂	CI	н
354 Z=CR+: V=CH: Y=0, X=N, n=2:	$R^{1}=$	CH ₃	OCH ₃	NH ₂	CI	Н

		- 1	ا	!	1	~· \		1
355	Z=CH; V=CH=CH; Y=CR ¹ ; X=N; n=1;	R ¹ =	CH ₃	OCH ₃	NH ₂	CI	н	
356	Z=CH: V=CH=CH: Y=CR ¹ : N=N: n=2;	R!=	CH ₃	OCH ₃	NH2	CI	Н	
357	Z=CH: V=N: Y=CH=C:R ¹); X=N: n=1:	R ^I =	CH ₃	OCH3	NH ₂	CI	Н	
358	Z=CH; V=N; Y=CH=C(R ¹); X=N; n=2;	R ^l =	CH ₃	осн ₃	NH ₂	Cl	н	1
359	$Z=CH; V=N; Y=C(R^{\frac{1}{4}})=N; X=CH; n=1;$	R1=	F	Cl	Br	CH ₃	н	
360	Z=CH; V=N; Y=C(R ¹ :=N; X=CH; n=2;	R1=	F	Cl	Br ˈ	CH ₃	н	
361	Z=CH: V=N: Y=CR ¹ : X=5: n=1:	R [{] =	F	CI.	Br	СНЗ	н	
362	$Z=CH: V=N: Y=CR^{T}: X=S: n=2:$	R1=	F	CI	Br	СНЗ	Н	
363	$Z=0; V=N; Y=CR^{T}; X=CH; n=1;$	R1=	F	CI	Br	CH ₃	н	
364	Z=O; V=N; Y=CR ¹ ; N=CH; n=2;	R 1 =	F	CI	Br	CH ₃	·H	
365	$Z=N; V=CH; Y=C(R^{\frac{1}{2}})=N; X=CH; n=1;$	R1=	F	CI	Br	СН3	Н.	
366	$Z=N_1 V=CH_1 Y=C(R^{\frac{1}{4}})=N_1 X=CH_1 n=2;$	R!=	F	CI	Вг	CH ₃	н	
367	Z=CH; V=N; Y=C(R:.=CH; X=CH; n=1;	R1-	F	C:	Br	CH ₃	н	
368	Z=CH: V=N: Y=C(R $^{\frac{1}{2}}$:=CH: X=CH: n=2:	R1=	F	CI	Br	CH ₃	Н	
369	Z=CH; V=CH; Y=NR ¹¹ ; X=N; n=1;	R ¹¹ =	CH ₃	C ₂ H ₅	Н	iPr	nPr	
370	Z=CH; V=CH; Y=NR ¹¹ ; X=N; n=2;	R11=	CH ₃	C ₂ H ₅	Н	iPr	nPr	1
371	Z=CH; V=N; Y=NR ¹¹ ; X=CH; n=1;	R11=	CH3	C ₂ H ₅	Н	iPr	nPr	
372	Z=CH; V=N; Y=NR ⁺¹ ; N=CH; n=2;	R11=	CH;	C ₂ H ₅	н	iPr	nPr	

Table 14

	·	. [CC	DLUMN		
	01=CG: R4=H	į	1	2	3	4	5
373	Z=S; N=CH; Y=CR ¹ ; N=N, n=1;	R1=.	CH ₃	осн;	NH ₂	CI	Н
374	Z=S: V=CH: Y=CR ¹ : N=N, n=2:	, R ¹ =	СН3	0СH3	NH2	Cl	н
375	Z=O; \'=CH; \'=CR\\. \\:N=N; n=1;	R-1=	CH ₃	осн;	NH ₂	CI	H
376	Z=O; V=CH; Y=CR ¹ , N=N, n=2;	R ! =	CH ₃	OCH ₃	NH ₂	Cl	н
377	Z=CH: V=S: Y=CR ¹ , N=N, n=1;	R1=	CH ₃	OCH ₃	NH2	CI	Н
378	Z=CH: N=S: Y=CR ¹ , N=N, n=2;	R 1 =	CH ₃	осн,	NH2	CI	Н
379	Z=CH; V=O; Y=CR ¹ , N=N, n=1;	R1=	СН3	осн ₃	NH ₂	CI	Н
380	Z=CH: V=O: Y=CR ¹ , N=N, n=2;	R1=	CH ₃	OCH ₃	NH ₂	CI	н
381	Z=CR ¹ : V=CH: Y=0, N=N, n=1:	R1=	CH3	OCH ₃	NH ₂	CI	н
382	Z=CR ¹ ; V=CH; Y=O, N=N, n=2;	R1=	CH3	OCH;	NH ₂	CI	Н
383	Z=CH: N=CH=CH: Y=CR ¹ : N=N: n=i:	R1=	CH3	OCH;	NH ₂	CI	Н
384	Z=CH, V=CH=CH; Y=CR † ; N=N; π =2,	R!=	· CH ₃	OCH3	NH ₂	CI	Н
385	Z=CH: N=N: Y=CH=C(R ¹): N=N: n=1:	R1=	CH;	осн3	NH ₂	CI	Н
385	Z=CH: V=N: Y=CH=C:R ¹ i: X=N: n=2:	R ! =	· CH3	OCH:	NH ₂	CI	H
387	Z=CH, V=N; Y=C(R ¹)=N; X=CH; n=1;	κ [:] =		CI	B7	CH ₃	Н

PCT/US94/08404

BEST AVAILABLE COPY

34

		:					
388	$Z=CH; N=N; Y=C(R^{T})=N; X=CH; n=2;$	R1=	F	CI	Br	СН₃	н
389	$Z=CH; V=N; Y=CR^{T}; X=S; n=1;$	R1=	F	CI	Br	СН3	н
390	Z=CH: V=N: Y=CR ¹ : X=S: n=2:	R1=	F .	CI	Br	CH ₃	Н
391	$Z=0; V=N; Y=CR^{T}; X=CH; n=1;$	R 1 =	F	CI	Br '	CH ₃	н
392	Z=O; V=N; Y=CR ¹ ; N=CH; n=2;	R!=	F	CI	Br	CH ₃	н
393	$Z=N; V=CH; Y=C(R^{T})=N; N=CH; n=1;$	R1=	F	CI	Br	CH ₃	н
394	$Z=N: V=CH: Y=C(R^{T})=N: X=CH: n=2:$	R !=	F	CI	Br	CH ₃	н
395	$Z=CH; V=N; Y=C(R^{T})=CH; X=CH; n=1;$	R ! =	F	CI	Br	CH ₃	н
396	$Z=CH; V=N; Y=C(R^{T})=CH; N=CH; n=2;$	R1=	F	CI	Br	СН3	н
397	Z=CH; V=CH; Y=NR ¹¹ ; X=N; n=1;	R11=	CH ₃	C ₂ H ₅	Н	iPr	nΩr
398	Z=CH: V=CH: Y=NR ¹¹ : N=N: n=2:	$R^{11} =$	CH;	C ₂ H ₅	Н	iPr	пРr
399	Z=CH; V=N; Y=NR ¹¹ ; X=CH; n=1;	R 11=	CH3	C2H5	Н	iPr	nPr
400	Z=CH; V=N; Y=NR ¹¹ ; X=CH; n=2;	R11=	CH:	C ₂ H ₅	н	iPr	nPr

Table 15

		ļ	COLUMN					
	$Q^1 = QG$: $R^4 = CH_3$	ļ	i	2	3	4	5	
401	Z=S; V=CH; Y=CR ¹ ; X=N, n=1;	R ! =	СН3	осн ₃	NH ₂	CI	Н	
402	Z=S; V=CH; Y=CR ¹ ; X=N, n=2;	R ! =	CH ₃	осн ₃	NH ₂	CI	Н	
403	Z=O; V=CH; Y=CR ¹ , X=N, n=1;	. R1=	CH ₃	осн ₃	NH ₂	Cl	Н	
404	Z=O; V=CH; Y=CR ¹ , X=N, n=2;	R1=	CH ₃	осн ₃	NH2	CI	H	
405	$Z=CH$; $V=S$; $Y=CR^{T}$, $X=N$, $n=1$;	$\mathbb{R}^1 =$	СН3	осн3	NH ₂	CI	н	
406	Z=CH: V=S: Y=CR ¹ , X=N, n=2:	R 1 =	CH ₃	осн ₃	NH ₂	Cı	н	
407	Z=CH; V=O; Y=CR ¹ , X=N, n=1;	R1=	CH ₃	осн ₃	NH ₂	CI	н	
408	Z=CH: V=O: Y=CR ¹ . N=N, n=2:	R ⁻¹ -=	CH3	ОСН3	NH ₂	-CI	H	-
409	Z=CR ¹ : V=CH: Y=O, X=N, n=1:	R1=	CH ₃	OCH ₃	NH ₂	Cı	н	
410	Z=CR ¹ ; V=CH; Y=O, X=N, n=2;	R1=	CH ₃	OCH ₃	NH ₂	CI	Н	
411	Z=CH; V=CH=CH; Y=CR ¹ ; X=N; n=1;	R1=	CH ₃	OCH ₃	NH ₂	CI	н	
412	Z=CH: V=CH=CH: Y=CR ¹ ; X=N: n=2:	R1=	CH3	осн ₃	NH ₂	CI	н	
413	$Z=CH$; $V=N$; $Y=CH=C(R^{\frac{1}{2}})$; $X=N$; $n=1$;	R1=	CH;	OCH ₃	NH ₂	Ci	Н	1
414	Z=CH; N=N; Y=CH=C+R ¹); X=N; n=2;	R1=	CH ₃	OCH;	NH ₂	CI	Н	į
415	$Z=CH; N=N; Y=C(R^{T})=N; X=CH; n=1;$	R 1 =	F	CI	Br	CH ₃	Н	1
416	Z=CH; N=N; Y=C(R ^T)=N; N=CH; n=2;	R 1 =	F	CI	Br	CH ₃	н	
417	$Z=CH; V=N; Y=CR^{T}; M=S; n=T;$	R 1 =	F	Cı	Br	CH3	Н	
418	Z=CH; N=N; Y=CR ¹ ; N=S; n=2.	R ! =	· : F	CI	Br	CH;	н	
419	$Z=0$; $N=N$; $Y=CR^{T}$; $N=CH$; $n=1$;	R!=	F	CI	Вг	CH ₃	H	
420	Z=0: V=N: Y=CR ¹ : X=CH: n=0:	R i =	F	CI	Br	CH ₃	н	:

WO 95/03306

BEST AVAILABLE COPY

35

421	Z=N; V=CH; Y=C(R ¹ =N; N=CH; π=1;	R ! =	F	Cl	Br	CH ₃	н	ı
422	$Z=N$; $V=CH$; $Y=C(R^{T})=N$; $X=CH$; $n=2$;	R1=	F	CI	Br	CH ₃	н	
423	Z=CH: V=N: Y=C(R) :=CH: N=CH: n=1;	R ^l =	F	CI	Br	CH ₃	н	
424	Z=CH: V=N: Y=C(R ¹ :=CH: N=CH: n=2:	R1=	F	CI	Br	CH ₃	н	
425	Z=CH; V=CH; Y=NR ¹¹ ; X=N; n=1;	R11=	CH ₃	C ₂ H ₅	н	iPr	nPr	
426	Z=CH: N=CH: Y=NR ¹⁴ ; N=N: n=2:	R11=	CH ₃	C2H5	Н	iPr	nPr	
427	Z=CH: V=N; Y=NR ¹¹ ; X=CH; n=1;	R11=	CH ₃	C ₂ H ₅	н	iPr .	nPr	
428	Z=CH; V=N; Y=NR ¹¹ ; X=CH; n=2;	R11=	CH;	C ₂ H ₅	Н	iPr	nPr	

Table 16

	.	. [COLUMN					
	<u>О¹=ОН</u>		i	. 2	3	4	5	ļ.
429	Z=S: $V=CH$: $Y=CR^{-1}$: $X=N$, $n=1$:	- R1=	СН3	осн ₃	ŃН ₂	CI	н	
430	Z=S: N=CH: Y=CR ¹ : N=N, n=2:	R 1 =	СНЗ	осн ₃	NH ₂	CI	Н	
431	Z=O; V=CH; Y=CR ¹ , X=N, n=1;	R1=	CH3	осн ₃	NH ₂	CI	Н	
432	Z=O; V=CH; Y=CR ¹ , X=N, n=2;	R ¹ =	CH ₃	осн3	NH ₂	CI	Н	
433	Z=CH: V=S: Y=CR ¹ , X=N, n=1;	R ¹ =	CH ₃	осн ₃	NH ₂	Cl	Н	
434	Z=CH; V=S; Y=CR ¹ , X=N, n=2;	R1=	CH ₃	осн ₃	NH2	Cl	н	
435	Z=CH; V=O; Y=CR ¹ , X=N, n=1;	R1=	CH ₃	OCH ₃	NH ₂	CI	н	
436	Z=CH; V=O; Y=CR ¹ , X=N, n=2;	R1=	CH ₃	осн ₃	NH ₂	CI	н	
437	Z=CR ¹ ; V=CH; Y=O, X=N, n=1;	R1=	CH ₃	осн3	NH ₂	CI	н	
438	Z=CR ¹ ; V=CH; Y=O, N=N, n=2;	R1=	CH3	OCH ₃	NH ₂	CI	н	
439	Z=CH: V=CH=CH: Y=CR ¹ ; X=N: n=1;	R1=	CH3	осн3	№Н2	Ci	H	
44()	Z=CH: V=CH=CH: Y=CR ¹ : X=N: n=2:	R1=	CH ₃	OCH ₃	NH ₂	CI	н -	1
441	Z=CH;-V=N;-Y=CH=C(R ⁺);-X=N;-n=1;-	R·l=	CH ₃	OCH3	NH2	CI	н	-
442	$Z=CH$; $V=N$; $Y=CH=C(R^{1})$; $N=N$; $n=2$;	R ! =	CH;	OCH ₃	NH ₂	CI	Н	
443	$Z=CH; V=N; Y=C(R^{\frac{1}{2}})=N; X=CH; n=1;$	R1=	F	CI	Br	CH3	Н	
1::	Z=CH; V=N; Y=C(R ¹ =N; X=CH; n=2;	R1=	F	CI	Br	CH ₃	Н	
445	Z=CH: N=N: Y=CR : X=S: n=1.	P. ! =	F	CI	Br	CH3	н	
446	$Z=CH(V=N)/Y=CR^{\frac{1}{2}}(X=S)/n=2)$	R1=	F	CI	Br	CH ₃	Н	١
447	Z=O: V=N: Y=CR ¹ : X=CH: n=1;	R!=	F	CI	Br	CH ₃	н	Ì
448	Z=O; V=N; Y=CR ¹ ; N=CH; n=2.	R1=	F	CI	Br	CH ₃	Н	
4-9	Z=N; V=CH; Y=C(R =N; X=CH; n=t;	R1=		CI	Br	CH ₃	н	
450	Z=N: V=CH: Y=C(R ¹ =N: N=CH: π=2:	R 1 =	F -	CI	Br	CH ₃	H	
451	Z=CH(V=N) Y=C(R) =CH, X=CH(n=1)	R [†] =	F	CI	Br	CH ₃	H	į
450	2 Z=CH; V=N; Y=C(R) =CH; N=CH; n=2.	R 1 =	: F	CI	Br	. CH ₃	Н	į
45	S Z=CH: V=CH: Y=NR ¹¹ ; X=N; n=1.	RII	= ! CH;	: C ₂ H ₅	н	iPr	nPr	į

Name of the Control o

PCT/US94/08404

36

454	Z=CH: V=CH: Y=NR ¹¹ : X=N: n=2:
455	Z=CH; V=N; Y=NR ¹¹ ; X=CH; n=1;
456	Z=CH; V=N; Y=NR ¹¹ ; N=CH; n=2;

R11=	СН3	C ₂ H ₅	н	iPr	nPr nPr
R11=	CH ₃	C ₂ H ₅	Н	iPr	nPr .
R ^{[[} =	CH;	C ₂ H ₅	Н	iPr	nPr

Table 17

		Į	COLUMN		<u>'</u>
	<u>0=01</u>		1	2	3
457	$Z=CH; V=N; Y=C(R^{T})=N; N=CH;$ R	1=	CI	Br	CH ₃
458	$Z=CH_1^{\frac{1}{2}}V=N_1^{\frac{1}{2}}Y=CR^{\frac{1}{2}}X=S_1^{\frac{1}{2}}$!=	CI	Br	СН3
459	$Z=0; V=N; Y=CR^{T}; N=CH;$ R	1=	CI	Br	CH ₃
460	$Z=N; V=CH; Y=C(R^{\frac{1}{4}})=N; X=CH;$ R	ξ ¹ =	Ci	Br	сн3
161	$Z=0: V=N: Y=CR^{-1}: X=N:$	۱ ₌	CI	Br	СН3
462	$Z=S; V=N; Y=CR^{-1}; X=N;$	= ا ۶	Cl	Br	СН3
463	$Z=CH; V=N; Y=NCH_3; X=CH;$	= ا ۶	Cl	Br	CH ₃

Table 18

		1	COLUMN		
	$\overline{O_1=O_1}$		ı	2	3
164	$Z=CH: V=N: Y=C(R^{T})=N: X=CH:$	R1=	Cl	Вг	СН3
465	$Z=CH$: $V=N$: $Y=CR^{T}$: $X=S$:	R1=	CI	Br	СН3
466	Z=0: V=N: Y=CR ¹ : X=CH:	R1=	CI	Br	CH ₃
467	$Z=N$; $V=CH$; $Y=C(R^{T})=N$; $X=CH$;	R 1 =	CI	Br	CH ₃
468	$Z=0; V=N; Y=CR^{-1}; X=N;$	$R^1 =$	Ci	Br	CH ₃
469	Z=S; V=N; Y=CR ¹ ; X=N;	R1=	CI	Br	CH ₃
470	Z=CH; V=N; Y=NCH ₃ ; X=CH;	$R^{l}=$	CI	Br	CH ₃

Table 19

	•		COLUMN				
	<u>0</u> !=0%		1	2	3	4	5
471	Z=0; V=N; Y=CR ¹ ; N=N; R 2 =H; n=1;	R1=	F	CI	Br	CH ₃	NH ₂
472	Z=0; V=N; Y=CR ¹ ; X=N; R ² =CH ₃ ; n=1.	R1=	F	Cl	Br	CH3	NH ₂
473	Z=0; $V=N$; $Y=CR^{\frac{1}{2}}$; $N=N$; $R^{\frac{1}{2}}=8n$; $n=1$;	R1=	F	CI	Br	CH3	NH ₂
474	Z=0; V=N; Y=CR ¹ ; X=N; R ⁴ =H; π =2;	R ! =	F	CI	Br	CH ₃	NH ₂
475	Z=O; V=N; Y=CR ¹ ; X=N, R ² =CH ₂ ; n=2;	R1=	F	Cı	Br	CH ₃	NH2
476	Z=O: V=N: Y=CR ¹ : X=N: R ² =Bn: n=2:	R1=	ŀ F	Cı	Br	CH3	SH ₂
477	$Z=S \cdot V=N \cdot Y=CR^{T} \cdot X=N \cdot R^{T}=H \cdot n=1$:	$R^{1}=$	·F.	CI	Br	CH:	NH.

		t t		1		1		
478	Z=S; V=N; Y=CR ¹ ; X=N; R ⁴ =CH ₃ ; n=1;	R ¹ =	F	CI	Br	CH ₃	NH ₂	
479	Z=S: $V=N$: $Y=CR^{-1}$: $N=N$: $R^{-2}=Bn$: $n=1$:	R ¹ =	F	CI	Br	CH ₃	NH2	
480	Z=S; $V=N$; $Y=CR^{-1}$; $X=N$; $R^{-4}=H$; $n=2$;	R ^l =	F	Cı	Br	CH ₃	NH ₂	
481	Z=S; $V=N$: $Y=CR^{-1}$: $X=N$: $R^{-2}=CH_{3}$: $n=2$:	R ^I =	F	CI	Br	CH ₃	NH2	
482	Z=S: $V=N$: $Y=CR^{T}$: $N=N$: $R^{A}=Bn$: $\pi=2$:	R1=	F	CI	Br	CH ₃	NH ₂	
483	$Z=N : V=N: Y=CR^{T}: X=0: R^{T}=H: \pi=1:$	R1=	F	CI	Br	CH ₃	NH ₂	
484	$Z=N: V=N: Y=CR^{+}: X=0: R^{+}=CH_{2}: n=1:$	R1=	F	Cl	Br	СН3	NH ₂	
485	$Z=N: V=N: Y=CR^{T}: X=O: R^{T}=Bn: \pi=1:$	R ^l =	F	CI	Br ·	сн3	NH ₂	
486	$Z=N: V=N: Y=CR^{T}: X=0: R^{4}=H: \pi=2:$	R1=	F	CI	Br	CH ₃	NH ₂	
487	$Z=N$; $V=N$; $Y=CR^{T}$, $X=O$; $R^{\frac{1}{2}}=CH_{3}$; $n=2$;	R1=	F	CI	Br	СН3	NH ₂	
488	Z=N : V=N: Y=CR ¹ : N=O: R ⁴ =Bn: n=2:	R1=	F	CI	Br	CH ₃	NH ₂	
489	$Z=CH: V=N: Y=C(R^{\frac{1}{4}})=N: X=CH: R^{\frac{2}{4}}=H: n=1;$	R ¹ =	CI	Br	CH ₃			
490	$Z=CH; V=N; Y=C(R^{\frac{1}{2}})=N; X=CH; R^{\frac{2}{4}}=H; n=2;$	R1=	Cl	Br	CH ₃			
491	Z=CH: V=N: Y=C(R $^{\frac{1}{2}}$ =N: X=CH: R $^{\frac{1}{2}}$ =CH ₃ : n=1:	R1=	CI	Br	CH ₃			
492	Z=CH, V=N; Y=C(R 1 =N; X=CH; R 4 =CH $_{3}$; n=2;	$\mathbb{R}^1 =$	CI	Br	CH ₃			
493	Z=CH: V=N: Y=CR ¹ : X=S: R ⁴ =H: n=1:	R1=	CI	Br	CH ₃			
191	Z=CH: V=N: Y=CR ¹ : X=S: R ⁴ =H: n=2:	R 1 =	CI	Br	CH ₃			
495	$Z=CH: V=N: Y=CR^{T}: X=S: R^{T}=CH_{3}: n=1;$	R 1=	CI	Вг	CH ₃			1
496	Z=CH; V=N; Y=CR ¹ ; X=S; R ⁴ =CH ₃ ; n=2;	$R^{1}=$	CI	Br	CH ₃			
497	Z=0: V=N: Y=CR ¹ : X=CH: R ⁴ =H: n=1;	$R^{l}=$	Cı	Br	CH ₃			
498	Z=0; V=N; Y=CR ¹ ; N=CH; R ⁴ =H; n=2;	R 1=	CI	Br	CH ₃			
499	Z=0: $V=N$: $Y=CR^{1}$: $N=CH$: $R^{4}=CH_{3}$: $n=1$:	R 1 =	Cı	Br	CH ₃			
500	Z=0: V=N: Y=CR ¹ : X=CH: R^4 =CH ₃ : n=2:	R1=	CI	Br	CH ₃			
501	$Z=CH, V=N; Y=NR^{+1}; X=CH; R^{+2}=H, n=1;$	R11=	CH ₃	Ει	iPr			
502	Z=CH: V=N: Y=NR ¹¹ ; N=CH; R ² =H, n=2;	R-1-1=	CH ₃	-Eı-	iPr—	ļ		-
503	Z=CH: V=N: Y=NR ¹¹ ; X=CH: R ⁴ =CH ₃ , n=1;	R ¹¹ =	CH ₃	Et	iPr	İ		1
504	$Z=CH: V=N: Y=NR^{1/2}: X=CH: R^2=CH_3, n=2;$	R11=	CH ₃	Ει	iPr			
505	$Z=N: V=CH: Y=C(R^{\frac{1}{2}})=N: X=CH: R^{\frac{1}{2}}=H: n=1;$	R1=	Cı	Br	CH ₃			
506	Z=N: V=CH: Y=C($R^{\frac{1}{4}}$ =N: X=CH: $R^{\frac{2}{4}}$ =H: n=2:	R1=	CI	Br	CH ₃	i		
507	Z=N: N=CH: Y=C(R $^{\frac{1}{2}}$ =N: N=CH: $R^{\frac{2}{2}}$ =CH ₃ : n=1:	R¹≓	CI	Br	CH ₃			
508	$Z=N, N=CH; Y=C(R^{\frac{1}{2}})=N; N=CH; R^{\frac{1}{2}}=CH_{\frac{1}{2}}; n=2.$	R1=	CI	Br	CH;			Ì

38

Table 20

		ĺ			COL	UMN	,
	<u> </u>	į	ı	2	3	4	5
509	Z=S: $V=CH$: $Y=CR^{-1}$: $X=N$. $R^{-4}=CH_{-3}$:	R1=	н	CI	CH ₃	осн ₃	O-n-hexyl
510	Z=S: $V=CH$: $Y=CR^{-1}$: $X=N$. $R^{-1}=H$:	R1=	Н	CI	CH ₃	осн ₃	O-n-hexyl
511	Z=O; V=CH; Y=CR ¹ ; N=N, R ⁴ =CH ₃ ;	R ^I =	Н	Cı	CH ₃	осн3	O-n-hexyl
512	Z=O: V=CH: Y=CR ¹ : N=N, R ⁴ =H;	R1=	Н	CI	СН3	осн ₃	O-n-hexyl
513	$Z=CR^{1}$; $V=CH$; $V=O$; $X=N$, $R^{4}=CH_{3}$;	R1= .	Н	CI .	СН3	осн3	O-n-hexyl
514	$Z=CR^{-1}$; $V=CH$; $Y=O$; $X=N$, $R^{-2}=H$;	R ¹ =	Н	CI	СН3	осн ₃	O-n-hexyl
515	Z=CR ¹ ; V=S; Y=CH; N=N, R ⁴ =CH ₃ ;	R ^I =	Н	CI	CH ₃	осн ₃	O-n-hexyl
516	$Z=CR^{-1}$; $V=S$; $Y=CH$; $X=N$, $R^{-4}=H$;	R¹=	Н	Cı	СН3	осн ₃	O-n-hexyl
517	Z=CR ¹ ; V=O; Y=CH; N=N, R ⁴ =CH ₃ ;	R 1 =	Н	CI	СН3	осн ₃	O-n-hexyl
518	Z=CR ¹ : V=O: Y=CH; N=N, R ⁴ =H;	R1=	Н	CI	CH ₃	осн3	O-n-hexyl
519	Z=CH: $V=S$: $Y=CR^{-1}$: $X=N$: $R^{-4}=CH_{-3}$:	R ^I =	Н	CI	СН₃	осн ₃	O-n-hexyl
520	$Z=CH; V=S; Y=CR^{1}; N=N, R^{4}=H;$	R ¹ =	н	Ci	СН3	осн ₃	O-n-hexyl
521	Z=CH: $V=0$: $Y=CR^{1}$: $X=N$: $R^{4}=CH_{3}$:	R ! =	н	CI	СН3	осн ₃	O-n-hexyl
522	$Z=CH: V=0: Y=CR^{1}: X=N, R^{4}=H;$	R 1 =	н	CI	СН3	осн ₃	O-n-hexyl
523	$Z=CR^{T}$; $V=CH=CH$; $Y=CH$; $X=N$, $R^{4}=CH_{3}$;	R ¹ =	Н	CI	CH ₃	осн ₃	O-n-hexyl
524	Z=CR ¹ : V=CH=CH; Y=CH; X=N, R ⁴ =H;	R ^I =	н	CI	CH ₃	осн ₃	O-n-hexyl
525	$Z=CR^{1}$: $V=CH=CH$: $Y=CH$: $X=N$. $R^{4}=CH_{3}$:	. R !=	н	CI	СН3	OCH ₃	O-n-hexyl
526	$Z=CR^{1}$; $V=N$; $Y=CH=CH$; $X=N$, $R^{4}=CH_{3}$;	R ¹ =	Н	. CI	CH ₃	осн ₃	O-n-hexyl
527	$Z=CR^{1}$; $V=N$; $Y=CH=CH$; $X=N$, $R^{4}=H$;	R ¹ =	н	CI	CH ₃	осн ₃	O-n-hexyl
528	Z=CH: V=CH=CH: Y=CR 1 ; X=N, R 4 =CH $_{3}$;	R 1 =	н	CI	CH ₃	осн ₃	O-n-hexyl
529-	Z=CH:-Y=CH=CH:-Y=CR ¹ : X=N: R ¹ =H:-	R-I =	- H-	-cı-	-CH3-	OCH ₃	O-n-hexyl
530	Z=CH; V=N; Y=CH=CR 1 ; X=N, R 4 =CH $_{3}$;	R1=	Н	CI	СН3	осн3	O-n-hexyl
531	Z=CH; V=N; Y=CH=CR; ; X=N, R ⁴ =H;	R ! =	Н	CI	CH ₃	осн ₃	O-n-hexyl
532	Z=CH: V=CH: $Y=NR^{11}$. $X=N$. $R^{4}=CH_{3}$:	R 11 =	н	Me	Εt	iPr	₁nPr
533	Z=CH: V=CH: Y=NR ¹¹ : N=N, R ⁴ =H:	R11=	Н	Me	Er	iPr	nPr

Table 21

			COLUMN					
	$O_1 = OVI$		<u> </u>	2	3_	1	5	
534	Z=CH, V=N; Y=CR =N; N=CH; n=1;	R¹=	F	CI	Br	СН3	н	
535	Z=CH: V=N: Y=CR ¹ =N: X=CH: n=2:	R ! =	F	CI	Br	CH ₃	н	
536	Z=CH; V=N; Y=CR ² ; N=S; n=1;	R !=	F	CI	Br	CH3	н	
537	Z=CH; N=N; Y=CR ² ; N=S; π=2;	R :=	F	CI	Br	CH ₃	Н і	

538	Z=0; V=N; Y=CR ¹ ; X=CH; n=1;	R1=	F	CI	Br	CH ₃	н	
539	Z=CH: V=N: Y=CR1: X=S: n=2:	R1=	F	CI	Br	CH ₃	н	
540	$Z=N; V=CH; Y=CR^{\frac{1}{2}}=N; X=N; n=1;$	R1=	F	CI	Br	CH ₃	н	
541	$Z=N; V=CH; Y=CR^{\frac{1}{2}}=N; X=N; n=2;.$	R1=	F	CI	Br	CH ₃	н	
542	Z=CH: V=N: Y=NR ¹¹ : X=CH: n=1:	R11=	н	CH ₃	C ₂ H ₅	iPr	nPr	
543	Z=CH; V=N; Y=NR ¹¹ ; X=CH; n=1;	R 11=	н	CH;	C ₂ H ₅	iPr	nPr	

Table 23

	Ī	COLUMN				
$Q^{\dagger} = QN R^{\frac{1}{2}} = H$	Ī	1	2	3	1	5
544 Z=S; V=CH; Y=CR ¹ ; X=N, n=1;	R!=	CH ₃	OCH ₃	NH2	Cl	н ·
545 Z=S: V=CH: Y=CR : X=N, n=2.	- R1=	сн3	осн _з	NH ₂	CI	н
546 Z=O: V=CH: Y=CR ¹ , N=N, n=1;	. R1=	СН3	OC43	NH ₂	CI	н
547 Z=O; N=CH; Y=CR [†] , N=N, n=2;	R ¹ =	CH ₃	OCH ₃	NH ₂	Cl	н
548 Z=CH: V=S: Y=CR ¹ , N=N, n=1:	R1=	СН3	осн ₃	NH ₂	CI	н
549 Z=CH: V=S: Y=CR ¹ , N=N, n=2;	R 1 =	CH ₃	осн ₃	NH ₂	Cı	н
550 Z=CH; V=O; Y=CR ¹ , X=N, n=1;	R 1 =	СН3	осн ₃	NH ₂	CI	н
551 Z=CH: V=O: Y=CR ¹ , N=N, n=2:	R 1 =	CH ₃	осн3	NH ₂	Cı	н
552 Z=CR ¹ ; V=CH; Y=O, X=N, n=1;	R 1 =	CH ₃	осн ₃	NH ₂	CI	н
553 Z=CR ¹ ; V=CH; Y=O, X=N, n=2;	R 1 =	CH ₃	OCH3	NH ₂	CI	н
554 Z=CH; V=CH=CH; Y=CR ¹ ; X=N; n=1;	R1=	CH ₃	OCH ₃	NH ₂	CI	н
555 Z=CH, V=CH=CH; Y=CR ^T ; N=N; n=2;	R ! =	CH ₃	OCH ₃	NH ₂	CI	Н
556 Z=CH; V=N; Y=CH=C(R ¹); X=N; n=1;	R1=	CH ₃	OCH ₃	NH ₂	CI	Н
557 Z=CH: N=N: Y=CH=C(R ¹); X=N: n=2;	R1=	CH ₃	OCH ₃	NH ₂	CI	,H
558 ZECHTVEN: YEC(R ¹)EN: NECHTAEI;	R-I =	- -F	CI	-Br	CH ₃	-H
559 Z=CH; N=N; Y=C(R ¹)=N; N=CH; n=2;	R1=	F	CI	Br	CH ₃	н
560 Z=CH: V=N: Y=CR1: N=S: n=1.	R!=	F	CI	Br	CH ₃	Н
561 Z=CH; N=N; Y=CR ¹ ; N=S; n=2.	R ! =	F	CI	Вг	CH ₃	н
562 Z=O; N=N; Y=CR ² ; N=CH; n=1;	R!=	F	Ci	Бг	CH ₃	Н
563 Z=O, N=N; Y=CR ³ ; N=CH; n=2.	R ! =	F	Ci	Br	CH ₃	Н
564 Z=N: V=CH: Y=C.R ¹ }=N: N=CH: n=1:	R1=	F	Ci	Вг	CH;	н
565 Z=N; V=CH; Y=C(R ¹)=N; N=CH; n=2;	81=	F	CI	Bř	CH3	Н
566 Z=CH: V=N: Y=C.R ¹ !=CH: X=CH: n=!;	R ! =	F	CI	Br	CH ₃	Ĥ
567 Z=CH; V=N; Y=C;R ³ ;=CH; N=CH; n=2;	R [!] =	F	CI	Br	CH;	ł:
508 Z=CH, V=CH; Y=NR ¹¹ ; N=N; n=1,	811	= : CH;	Calls	H	i iPr	n?r
569 Z=CH; V=CH; Y=NR ¹⁴ ; N=N; n=2;	RU	= CH;	Calis	H	- iPr	nPr
570 Z=CH, N=N, Y=NR ¹¹ , N=CH; n=1;	R!!	= : CH	C ₂ H ₅	н	, iPr	nl'r

571 Z=CH: V=N: Y=NR¹¹: X=CH: n=2:

Table 23

			COLUMN				
	$Q^{\dagger} = QN R^{\frac{1}{2}} = CH_3$		1	2	3	4	5
572	Z=S: V=CH; Y=CR ¹ ; X=N, n=!;	R1=	CH ₃	осн ₃	NH ₂	CI	н
573	Z=S; V=CH; Y=CR ¹ ; N=N, n=2;	R!=	СН3	OCH ₃	NH ₂	CI	н
574	Z=0: Y=CH: Y=CR ¹ , X=N, n=1:	R¹=	CH ₃	осн3	NH ₂	Cl	н
575	Z=0: V=CH: Y=CR ¹ , N=N, n=2:	RI=	CH_3	осн ₃	NH2	CI	н
576	Z=CH: V=S; Y=CR ¹ , X=N, n=1;	R1=	CH ₃	осн ₃	NH ₂	C1	Н
577	Z=CH; V=S; Y=CR ¹ , X=N, n=2;	R ⁱ =	CH ₃	осн ₃	NH ₂	Cl	н
578	Z=CH; V=O; Y=CR ¹ , N=N, n=1;	R1=	CH ₃	осн ₃	NH ₂	Cl	н
579	Z=CH: V=O: Y=CR ¹ , X=N, n=2;	R 1 =	CH ₃	осн ₃	NH ₂	CI	н
580	Z=CR ¹ ; V=CH; Y=O, N=N, n=1;	R ! =	CH ₃	осн ₃	NH2	CI	н
581	Z=CR ¹ ; V=CH; Y=O, X=N, n=2;	R 1 =	сн3	осн ₃	NH ₂	CI	н.
582	Z=CH; V=CH=CH; Y=CR ¹ ; X=N; n=1;	R 1 =	CH ₃	OCH ₃	NH ₂	Ci	н
583	Z=CH; V=CH=CH; Y=CR ¹ ; X=N; n=2;	R1=	CH ₃	осн ₃	NH ₂	CI	н
584	$Z=CH; V=N; Y=CH=C(R^{1}); X=N; n=1;$	R!=	CH ₃	осн3	NH ₂	CI	Н
585	$Z=CH$; $V=N$; $Y=CH=C(R^{1})$; $N=N$; $n=2$;	R1=	CH ₃	OCH ₃	NH ₂	CI	н
586	$Z=CH; V=N; Y=C(R^{T})=N; X=CH; n=1;$	R!=	F	CI	Br	CH ₃	Н
587	$Z=CH$: $V=N$: $Y=C(R^{1})=N$: $X=CH$: $n=2$;	R1=	F	CI	Br	CH ₃	Н
588	$Z=CH; V=N; Y=CR^{1}; N=S; n=1;$	R1=	F	CI	Br	CH ₃	н
589	$Z=CH: V=N: Y=CR^{T}: N=S: n=2:$	R ¹ =	F	CI	Br	CH ₃	Н
590	Z=O: V=N: Y=CR ¹ : N=CH: n=1:	R1=	F	CI	Br	CH ₃	н
591	Z=O: V=N: Y=CR ¹ : X=CH: n=2:	RI=	F	CI	Br	CH ₃	Н
592	$Z=N; V=CH; Y=C(R^{T})=N; X=CH; n=1;$	R1=	F	CI	Br	CH ₃	,H
593	$Z=N; V=CH; Y=C(R^{\frac{1}{2}})=N; X=CH; n=2;$	R ! =	F	CI	Br	CH ₃	н
594	Z=CH; V=N; Y=C($\mathbb{R}^{\frac{1}{2}}$;=CH; X=CH; n=1;	R ¹ =	F	CI	Br	CH ₃	н
595	$Z=CH$; $Y=N$; $Y=C(R^T)=CH$; $X=CH$; $n=2$;	R 1 =	F	CI	Br	CH ₃	н
596	S Z=CH: V=CH: Y=NR ¹¹ ; X=N: n=1:	R11=	E CH	C ₂ H ₅	н	iPr	nPr
59	7 Z=CH: V=CH: Y=NR ¹¹ ; X=N: n=2:	* R ¹¹ :	- CH:	C ₂ H ₅	н	iPr	nPr
59	S Z=CH: V=N: Y=NR ¹¹ : X=CH: n=1:	R11:	= CH	C_2H_5	Н	iPr	nPr
59	Z=CH; V=N; Y=NR ¹¹ ; N=CH; n=2;	R^{11}	= <u> CH</u>	CoH5	н	iPr	nPr

Formulation/Utility

Compounds of this invention will generally be used in formulation with an agriculturally suitable carrier comprising a liquid or solid diluent. Useful formulations

powders, emulsifiable concentrates, dry flowables and the like, consistent with the physical properties of the active ingredient, mode of application and environmental factors such as soil type, moisture and temperature. Sprayable formulations can be extended in suitable media and used at spray volumes from about one to several hundred liters per hectare. High strength compositions are primarily used as intermediates for further formulation. The formulations will typically contain effective amounts of active ingredient, diluent and surfactant within the following approximate ranges which add up to 100 weight percent.

	Weight Percent		
	<u>Active</u> Ingredient	Diluent	Surfactant
Wettable Powders	5-90	0-74	1-10
Oil Suspensions, Emulsions, Solutions, (including Emulsifiable Concentrates)	5-50	40-95	0-15
Dusts	1-25	70-99	0-5
Granules, Baits and Pellets	0.01-99	5-99.99	0-15
High Strength Compositions	90-99	0-10	0-2

10

15

20

25

Typical solid diluents are described in Watkins, et al., Handbook of Insecticide Dus: Diluents and Carriers, 2nd Ed., Dorland Books, Caldwell, New Jersey: Typical liquid diluents and solvents are described in Marsden, Solvents Guide, 2nd Ed.,

Interscience, New York, 1950. McCutcheon's Detergents and Emulsifiers Annual, Allured Publ. Corp., Ridgewood, New Jersey, as well as Sisely and Wood. Encyclopedia of Surface Active Agents, Chemical Publ. Co., Inc., New York, 1964, list surfactants and recommended uses. All formulations can contain minor amounts of additives to reduce foam, caking, corrosion, microbiological growth, and the like.

Solutions are prepared by simply mixing the ingredients. Fine solid compositions are made by blending and, usually, grinding as in a hammer mill or fluid energy mill. Water-dispersible granules can be produced by agglomerating a fine powder composition; see for example, Cross et al., *Pesticide Formulations*, Washington, D.C., 1988, pp 251-259. Suspensions are prepared by wet-milling; see, for example, U.S. 3,060,084. Granules and pellets can be made by spraying the active material upon preformed granular carriers or by agglomeration techniques. See Browning. "Agglomeration", *Chemical Engineering*, December 4, 1967, pp 147-148, *Perry's*

WO 95/03306

10

BEST AVAILABLE COPY

42

Chemical Engineer's Handbook, 4th ed., McGraw-Hill, New York, (1963), pp 8-57 and following, and WO 91/13546.

For further information regarding the art of formulation, see U.S. 3,235,361, Col. 6, line 16 through Col. 7, line 19 and Examples 10-41; U.S. 3,309,192, Col. 5, line 43 through Col. 7, line 62 and Examples 8, 12, 15, 39, 41, 52, 53, 58, 132, 138-140, 162-164, 166, 167 and 169-182; U.S. 2,891,855, Col. 3, line 66 through Col. 5, line 17 and Examples 1-4; Klingman, Weed Control as a Science, John Wiley and Sons, Inc., New York, (1961), pp 81-96; and Hance et al., Weed Control Handbook, 8th ed., Blackwell Scientific Publications, Oxford, (1989).

In the following Examples, all percentages are by weight and all formulations are prepared in conventional ways. Compound numbers refer to compounds in Index Table A.

F	c n	יוח	nΙ	و	Ā

		•
	Wettable Powder	
15	Compound I	65.0%
	dodecylphenol polyethylene glycol ether	2.0%
	sodium ligninsulfonate	4.0%
	sodium silicoaluminate	6.0%
	montmorillonite (calcined)	23.0%.
20	Example B	
	Granule	
	Compound 1	10.0%
	attapulgite granules (low volatile	
	matter, 0.71/0.30 mm; U.S.S. No.	
25	25-50 sieves)	90.0%.
	Example C	
	Extruded Pellet	
	Compound 1	25.0%
	anhydrous sodium sulfate	10.0%
30	crude calcium ligninsulfonate	5.0%
	sodium alkylnaphthalenesulfonate	1.0%
	calcium/magnesium bentonite	59.0%.
	Example D	
	Emulsifiable Concentrate	
. 35	Compound :	20.0₹
	plend of oil soluble sulfonates	
	and polyoxyethylene ethers	10.0%
	isophorone	70.0℃.

Commission of the contract of the contract of

The compounds of this invention exhibit activity against a wide spectrum of foliar-feeding, fruit-feeding, stem or root feeding, seed-feeding, aquatic and soil-inhabiting arthropods (term "arthropods" includes insects, mites and nematodes) which are pests of growing and stored agronomic crops, forestry, greenhouse crops, ornamentals, nursery crops, stored food and fiber products, livestock, household, and public and animal health. Those skilled in the art will appreciate that not all compounds are equally effective against all growth stages of all pests. Nevertheless, all of the compounds of this invention display activity against pests that include: eggs, larvae and adults of the Order Lepidoptera; eggs, foliar-feeding, fruit-feeding, root-feeding, seed-feeding larvae and adults of the Order Coleoptera; eggs, immatures 10 and adults of the Orders Hemiptera and Homoptera; eggs, larvae, nymphs and adults of the Order Acari; eggs, immatures and adults of the Orders Thysanoptera, Orthoptera and Dermaptera; eggs, immatures and adults of the Order Diptera; and eggs, junveniles and adults of the Phylum Nematoda. The compounds of this invention are also active against pests of the Orders Hymenoptera, Isoptera, Siphonaptera, Blattaria, Thysanura 15 and Psocoptera; pests belonging to the Class Arachnida and Phylum Platyhelminthes. Specifically, the compounds are active against southern corn rootworm (Diabrotica undecimpunctata howardi), aster leafhopper (Mascrosteles fascifrons), boll weevil (Anthonomus grandis), two-spotted spider mite (Tetranychus urticae), fall armyworm (Spodoptera frugiperda), black bean aphid (Aphis fabac), green peach aphid (Myzus 20 persica), cotton aphid (Aphis gossypii), Russian wheat aphid (Diuraphis noxia), English grain aphid (Sitobion avenue), tobacco budworm (Heliothis virescens), rice water weevil (Lissorhoptrus oryzophilus), rice leaf beetle (Oulema oryzae), whitebacked planthopper (Sogatella furcifera), green leafhopper (Nephotettix cincticeps), brown planthopper (Nilaparvata lugens), small brown planthopper (Laodelphax striatellus), 25 rice stem borer (Chilo suppressalis), rice leafroller (Cnaphalocrocis medinalis), black rice stink bug (Scotinophara lurida), rice stink bug (Oebalus pugnax), rice bug (Leptocorisa chinensis), slender rice bug (Cletus pantiger), and southern green stink bug (Nezara viridada). The compounds are active on mites, demonstrating ovicidal, larvioldal and chemosterilant activity against such families as Tetranychidae including 30 Tetranychus urucae, Tetranychus cinnabarinus, Tetranychus medanieli, Tetranychus paziticus, Tetranychus turkestani, Byrobia rubrioculus, Panonychus ulmi, Panonychus ciri, Eotetranychus carpini borcalis, Eotetranychus, hicoriae, Eotetranychus sexumentatus, Entetranycius yumensis, Entetranycius banksi and Oligonychus 35 printensist. Tennicalvidae including Brevipalpus iewisi, Brevipalpus phoenicis, Brevipalpus californicus una Brevipalpus obovatus. Eriophyidae including Phyllocopurum oleivara. Eriophyes sheldoni, Aculus cornuus. Epitrimerus pyri and

WO 95/03306

5

10

15

20

30

35

PCT/US94/08404

44

Eriophyes mangiferne. See WO 90/10623 and WO 92/00673 for more detailed pest descriptions.

Compounds of this invention can also be mixed with one or more other insecticides, fungicides, nematocides, bactericides, acaricides, growth regulators, chemosterilants, semiochemicals, repellants, attractants, pheromones, feeding stimulants or other biologically active compounds to form a multi-component pesticide giving an even broader spectrum of agricultural protection. Examples of other agricultural protectants with which compounds of this invention can be formulated are: insecticides such as avermeetin B, monocrotophos, carbofuran, tetrachlorvinphos, malathion, parathion-methyl, methomyl, chlordimeform, diazinon, deltamethrin, oxamyl, fenvalerate, esfenvalerate, permethrin, profenofos, sulprofos, triflumuron. diflubenzuron, methoprene, buprofezin, thiodicarb, acephate, azinphosmethyl, chlorpyrifos, dimethoate, fipronil, flufenprox, fonophos, isofenphos, methidathion, metha-midophos, phosmet, phosphamidon, phosalone, pirimicaro, phorate, terbufos, trichlorfon, methoxychlor, bifenthrin, biphenate, cyfluthrin, tefluthrin, fenpropathrin, fluvalinate, flucythrinate, tralomethrin, imidacloprid, metaldehyde and rotenone; fungicides such as carbendazim, thiuram, dodine, maneb, chloroneb, benomyl, cymoxanil, fenpropidine, fenpropimorph, triadimefon, captan, thiophanate-methyl, thiabendazole, phosethyl-Al, chlorothalonil, dichloran, metalaxyl, captafol, iprodione, oxadixyl, vinclozolin, kasugamycin, myclobutanil, tebuconazole, difenoconazole, diniconazole, fluquinconazole, ipconazole, metconazole, penconazole, propiconazole, uniconzole, flutriaiol, prochloraz, pyrifenox, fenarimol, triadimenol, diclobutrazol, copper oxychloride, furalaxyl, folpet, flusilazol, blasticidin S, diclomezine, edifenphos, isoprothiolane, iprobenfos, mepronil, neo-asozin, pencycuron, probenazole, pyroquilon, tricyclazole, validamycin, and flutolanil; nematocides such as aldoxycarb, fenamiphos 25 and fosthietan; bactericides such as oxytetracyline, streptomycin and tribasic copper sulfate; acaricides such as binapacryl, oxythioquinox, chlorobenzilate, dicofol, dienochlor, cyhexatin, hexythiazox, amitraz, propargite, tebufenpyrad and fenbutatin oxide; and biological agents such as entomopathogenic bacteria, virus and fungi.

In certain instances, combinations with other arthropodicides having a similiar spectrum of control but a different mode of action will be particularly advantageous for resistance management.

Arthropod pests are controlled and protection of agronomic, horticultural and specialty crops, animal and human health is achieved by applying one or more of the compounds of this invention, in an effective amount, to the environment of the pests including the agrenomic and/or nonagronomic locus of infestation, to the area to be protected, or directly on the pests to be controlled. Thus, the present invention further comprises a method for the control of foliar and soil inhabiting arthropods and

30

45

applying one or more of the compounds of Formula I or Formula II, or compositions containing at least one such compound, in an effective amount, to the environment of the pests including the agronomic and/or nonagronomic locus of infestation, to the area to be protected, or directly on the pests to be controlled. A preferred method of application is by spraying. Alternatively, granular formulations of these compounds can be applied to the plant foliage or the soil. Other methods of application include direct and residual sprays, aerial sprays, seed coats, microencapsulations, systemic uptake, baits, eartags, boluses, foggers, fumigants, aerosols, dusts and many others. The compounds can be incorporated into baits that are consumed by the arthropods or in devices such as traps and the like.

The compounds of this invention can be applied in their pure state, but most often application will be of a formulation comprising one or more compounds with suitable carriers, diluents, and surfactants and possibly in combination with a food depending on the contemplated end use. A preferred method of application involves spraying a water dispersion or refined oil solution of the compounds. Combinations with spray oils, spray oil concentrations, spreader stickers, adjuvants, and synergists and other solvents such as piperonyl butoxide often enhance compound efficacy.

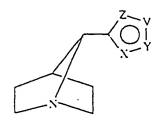
The rate of application required for effective control will depend on such factors as the species of arthropod to be controlled, the pest's life cycle, life stage, its size, location, time of year, host crop or animal, feeding behavior, mating behavior, ambient moisture, temperature, and the like. Under normal circumstances, application rates of about 0.01 to 2 kg of active ingredient per hectare are sufficient to control pests in agronomic ecosystems, but as little as 0.001 kg/hectare may be sufficient or as much as 8 kg hectare may be required. For nonagronomic applications, effective use rates will range from about 1.0 to 50 mg/square meter but as little as 0.1 mg/square meter may be sufficient or as much as 150 mg/square meter may be required.

The following TESTS demonstrate the control efficacy of compounds of this invention on specific pests. "Control efficacy" represents inhibition of arthropod development finelading mortality) that causes significantly reduced feeding. The pest control protection afforded by the compounds is not limited, however, to these species. See Index Tables A-D for compound descriptions.

BEST AVAILABLE COPY

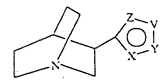
PCT/US94/08404

46 Index Tuble A



Compound	<u>Z</u>	$\overline{\lambda}$	<u>Y</u>	\mathbf{X}	<u>m.p. °C</u>
1	СН	N	C(CI)=CH	СН	77-78
2	СН	N	CH=CH	· CH	. oil ^a
3	СН	N	C(C1)=C(C1)	CH	30-32
4	N	СН	$C(CH_3)=N$	СН	oil ^b

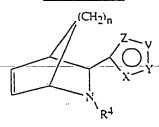
Index Table B



Compound
5

Z Y Y X m.p. °C N CH CH=N CH 45-50

Index Table C



Compound	<u>Z</u>	7.	$\underline{\mathbf{Y}}$	X	<u>R</u> ±	<u>n</u>	<u>m.p. -</u> C
6	0	N	C(CH ₃)	N	Bn	l	oii
7	CH	N.	CH=CH	CH	CH ₂	2	b_{lio}

47 Index Table D

			J		
Compound	. <u>7</u>	<u>Y</u> .	<u>. X</u> .	R4	<u>m.p. °C</u>
S	СН	C(CI)=CH	CH	H	159-161

¹H NMR (CDCI₃ DATA)

δ 8.65 (s.1H), 8.50 (d,1H), 7.80 (d,1H), 7.25 (d,1H), 3.81 (s.1H), 3.10-3.00 5 (m,1H), 3.00-2.95 (m,1H), 2.80-2.60 (m,2H), 2.50-2.40 (m,1H), 1.90-1.80 (m,1H), 1.45-1.10 (m,3H).

δ 8.68 (s.1H), 8.39 (s.1H), 3.87 (s.1H), 3.2-3.1 (m,2H), 2.85-2.65 (m,2H), 2.55 (s.3H), 2.6-2.5 (m,1H), 1.9-1.8 (m,1H), 1.5-1.2 (m,3H).

δ 7.35-7.2 (m.5H), 6.6 (dd.1H), 6.4 (dd.1H), 3.97 (s.1H), 3.6 (ABq.2H), 3.2 10 (s,1H), 3.0 (s,1H), 2.34 (s,3H), 2.05 (d,1H), 1.55 (d,1H).

\$ 8.66 (s.1H), \$.45 (d.1H), 7.8 (d.1H), 7.8 (d.1H), 7.25-7.20 (m.1H), 6.6 (dd.1H). ċ 6.4 (dd.1H), 3.45-3.40 (m.1H), 3.0 (s.1H), 2.45-2.40 (m.1H), 2.17 (s.3H), 2.05-1.95 (m.1H), 1.4-1.2 (m.2H), 0.95-0.80 (m.1H).

15

20

TEST A

Fall Armyworm

Test units, each consisting of a H.I.S. (high impact styrene) tray with 16 cells were prepared. Wet filter paper and approximately 8 cm² of lima bean leaf was placed into twelve of the cells. A 0.5 cm layer of wheat germ diet was placed into the four remaining cells. Fifteen to twenty third-instar larvae of fall armyworm (Spodoptera frugiperda) were placed into an 8 ounce (230 mL) plastic cup. Solutions of each of the test compounds in 75/25 acetone/distilled water solvent were sprayed into the tray and cup. Spraying was accomplished by passing the tray and cup, on a conveyer belt, directly beneath a flat fan hydraulic nozzle which discharged the spray at a rate of 0.5 bounds of active ingredient per acre (about 0.55 kg/ha) at 30 p.s.i. (207 kPa). The insects were transferred from the Sounce cup into the cells of the H.I.S. tray (one insect

per cell). The travs were covered and held at 27°C and 50% relative humidity for 48 h.

WO 95/03306

5

10

15

20

25

30

BEST AVAILABLE COPY

PCT/US94/08404

48

after which time readings were taken on the twelve cells with lima bean leaves. The four remaining cells were read 7 days later for delayed toxicity readings. Of the compounds tested, the following gave control efficacy levels of 80% or higher: 3*.

TEST B

Southern Corn Rootworm

* - tested at 250 ppm.

Test units, each consisting of an 8 ounce (230 mL) plastic cup containing a one-inch square (2.54 cm²) of a wheatgerm diet, were prepared. Solutions of each of the test compounds in 75/25 acetone/distilled water solvent were sprayed into the tray and cup. Spraying was accomplished by passing the tray and cup, on a conveyer belt, directly beneath a flat fan hydraulic nozzle which discharged the spray at a rate of 0.5 pounds of active ingredient per acre (about 0.55 kg/ha) at 30 p.s.i. (207 kPa). After the spray on the cups had dried, five second-instar larvae of the southern corn rootworm (Diabrotica undecimpurctata howardi) were placed into each cup. The cups were then held at 27°C and 50% relative humidity for 48 h. after which time mortality readings were taken. The same units were read again at 8 days. Of the compounds tested, the following gave control efficacy levels of 80% or higher: 1, 3* and 6.

* Test conducted at 250 ppm.

TEST C

Aster Leafhopper

Test units were prepared from a series of 12 ounce (350 mL) cups, each containing out (Avena sativa) seedlings in a 1 inch (2.54 cm) layer of sterilized soil. The test units were sprayed as described in TEST A with individual solutions of the test compounds. After the outs had dried from the spraying, between 10 and 15 adult aster leathoppers (Mascrosteles fascifrons) were aspirated into each of the cups. The cups were covered with vented lids and held at 27°C and 50% relative humidity for 48 h, after which time mortality readings were taken. Of the compounds tested, the following gave mortality levels of 80% or higher: 1*, 3*.

* Test conducted at 250 ppm.

49

TEST D

Two-Spotted Spider Mite

One inch squares (2.54 cm) of kidney bean leaves that had been infested on the undersides with 25 to 30 adult mites (*Tetranychus urticae*) were sprayed with their undersides facing up on a hydraulic sprayer with a solution of the test compound in 75/25 acetone/distilled water solvent. Spraying was accomplished by passing the leaves, on a conveyor belt, directly beneath a flat fan hydraulic nozzle which discharged the spray at a rate of 0.55 pounds of active ingredient per acre (about 0.5 kg/ha) at 30 p.s.i. (207 kPa). The leaf squares were then placed underside-up on square of wet cotton in a petri dish and the perimeter of the leaf square was tamped down onto the cotton with forceps so that the mites cannot escape onto the untreated leaf surface. The test units were held at 27°C and 50% relative humidity for 48 h, after which time mortality readings were taken. Of the compounds tested, the following gave mortality levels of 80% or higher: 1, 5 and 6.

15

20

30

10

TEST E

Boll Weevil

Test units consisting of 9 ounce (260 mL) cups containing five adult boll weevils (Anthonomus grandis grandis) were prepared. The test units were sprayed as described in TEST A with individual solutions of the test compounds. Each cup was covered with a vented lid and held at 27°C and 50% relative humidity for 48 h, after which time mortality readings were taken. Of the compounds tested, the following gave mortality levels of 80% or higher: 1, 2°.

* Test conducted at 250 ppm

TEST F

25 Contact Test Against Black Bean Aphid

Individual nasturtium leaves were infested with 10 to 15 aphids (all morphs and growth stages of *Aphis fabac*) and sprayed with their undersides facing up as described in TEST A. The leaves were then set in 3/8 inch (0.94 cm) diameter vials containing 4 mL of sugar solution (approximately 1.4 g per liter) and covered with a clear plastic 1 cunce (29 mL) cup to prevent escape of the aphids that drop from the leaves. The test units were held at 27°C and 50% relative humidity for 48 h, after which time mortality readings were taken. Of the compounds tested, the following gave mortality levels of 80% or higher: 1, 2*, 3*.

CLAIMS

A compound of the formula 1.

$$\mathbb{R}^{1} \longrightarrow \mathbb{R}^{2}$$

5

wherein:

Q is selected from the group

 $(CH_2)_n$

Q-1

Q-2

and

$$(CH_2)_n$$

$$(CH_2)_q$$

$$(R^3)_0$$

Q-3

Q-4

10

15

where the broken line represents an optional chemical bond;

 R^1 and R^2 are independently selected from the group H, halogen, C_1 - C_6 alkyl. C1-C6 haloalkyl. C2-C6 alkenyl, C2-C6 haloalkenyl, C2-C6 alkynyl, C2-C6 haloalkynyl, C3-C6 cycloalkyl, C3-C6 halocycloalkyl, CN, SCN, NO2, $N(R^5)R^6$, OR^5 , $C(O)R^5$, $C(O)OR^5$, $C(O)N(R^5)R^6$, SR^5 , $S(O)R^5$, $S(O)_7R^5$, S(O)₂N(R⁵)R⁶ and C₁-C₆ alkyl substituted with 1 or 2 groups independently selected from NO2, CN, C1-C3 alkylthio, C1-C3 alkoxy. C₁-C₃ haloalkoxy, C₂-C₄ alkylearbonyl and C₂-C₄ alkoxycarbonyl; R² being attached to any unsubstituted aromatic ring carbon; and R1 and R2 are not both hydrogen when Q is Q-1 or Q-4, n is 1, R³ is H and q is 2; 20

51

	R ³ , which is attached to any carbon of the azableyetic ring including the carbon
	directly attached to the heterocyclic aromatic ring, is selected from the
	group H. halogen. C1-C6 alkyl, C1-C6 haloalkyl, C2-C6 alkenyl, C2-C6
	haloalkenyl, C2-C6 alkynyl, C2-C6 haloalkynyl, C3-C6 cycloalkyl, C3-C6
5	halocycloalkyl, CN, SCN, NO ₂ , N(R ⁷)R ⁸ , OR ⁷ , C(O)R ⁷ , C(O)OR ⁷ ,
	$C(O_1N(R^7)R^8, SR^7, S(O)R^7, S(O)_2R^7, S(O)_2N(R^7)R^8 \text{ and } C_1-C_6 \text{ alkyl}$
	substituted with 1 or 2 groups independently selected from NO2, CN,
	C_1 - C_3 alkylthio, C_1 - C_3 alkoxy, C_1 - C_3 haloalkoxy, C_2 - C_4 alkylcarbonyl and
	C ₂ -C ₂ alkoxycarbonyl;
10	R^4 is selected from the group H, C_1 - C_6 alkyl, C_3 - C_6 cycloalkyl, C_1 - C_6 haloalkyl, C_2 - C_6 alkenyl, C_3 - C_6 alkynyl, $N(R^9)R^{10}$, $C(0)R^9$, $C(0)OR^9$,
	$C(O)N(R^9)R^{10}$, SR^9 , $S(O)R^9$, $S(O)_2R^9$, $S(O)_2N(R^9)R^{10}$, benzyl and
	CHTCH ₃ :Ph: provided when any of R ¹ , R ² , R ³ or R ⁴ is $S(O)R^5$, $S(O)_2R^5$.
	$S(O)R^7$, $S(O)_2R^7$, $S(O)R^9$, or $S(O)_2R^9$ then R^5 , R^7 and R^9 are other than H;
15	R ⁵ , R ⁶ , R ⁷ , R ⁸ , R ⁹ and R ¹⁰ are independently selected from the group H, C ₁ -C ₆
	alkyl, C ₁ -C ₆ haloalkyl, C ₃ -C ₆ cycloalkyl, phenyl optionally substituted with
	I or 2 substituents independently selected from W, and benzyl optionally
	substituted with 1 or 2 substitutents independently selected from W;
	W is selected from the group halogen, NO ₂ , CN, C ₁ -C ₃ alkyl, C ₁ -C ₃ haloalkyl,
20	C_1 - C_3 alkylthio, C_1 - C_3 alkoxy, C_1 - C_3 haloalkoxy, C_2 - C_4 alkylcarbonyl and
	C ₂ -C ₄ alkoxycurbonyl;
	m and n are independently 0, 1 or 2;
	p is 1 or 2; and
	q is 1, 2 or 3.

25

- 2. A compound according to Claim 1 wherein Q is Q-1.
- 3. A compound according to Claim 2 wherein

 R¹ is selected from the group H, halogen and C₁-C₂ alkyl;

 S² is selected from the group H and Cl;

 R³ is selected from the group H, halogen, C₁-C₆ alkyl and OR⁷;

 R⁵ is selected from the group H and C₁-C₄ alkyl; and a is 0 or 1.
- 35 4. A compound according to Claim 3 which is

7-(6-chloro-3-pyridinyl)-1-azabicyclo[2,2,1]heptane.

5. A method for controlling arthropods comprising contacting the arthropods or their environment with an arthropodicidally effective amount of a compound of the formula:

5

10

П

wherein:

Q1 is selected from the group

$$(R^{3})_{p} = (CH_{2})_{n}$$

$$(CH_{2})_{n}$$

$$(CH_{2})_{p}$$

$$(CH_{2})_{q}$$
and
$$(CH_{2})_{m}$$

$$(R^{3})_{p}$$

$$(R^{3})_{p}$$

$$(CH_{2})_{q}$$

$$(R^{3})_{p}$$

$$(R^{3})_{p}$$

$$(R^{3})_{p}$$

$$(R^{3})_{p}$$

$$(R^{3})_{p}$$

$$(R^{3})_{p}$$

where the broken line represents an optional chemical bond;

V, X, Y and Z of the ring are each independently selected from the group O, S, N, $-C(R^1)$ -, $-C(R^1)=C(R^2)$ -, $-C(R^1)=N$ - and $-N(R^{11})$ -; provided that (i) no more than one of V, X, Y or Z is $-C(R^1)=C(R^2)-, -C(R^1)=N-, -N(R^{11})-, O$ 15 or S. (ii) at least one of V, X, Y or Z is N, (iii) when the ring is a 5membered ring containing two N and one O or S, and Q is Q-6, then the ring is attached to the 2-position of Q1 and (iv) when the ring is a 5membered ring containing two N and one O or S, then Q is other than Q-5; R^1 and R^2 are independently selected from the group H, halogen, C_1 - C_6 alkyl, 20 C1-C5 haloalkyl, C2-C6 alkenyl, C2-C6 haloalkenyl, C2-C6 alkynyl, C2-C6 haloalkynyl: C3-C6 cycloalkyl, C3-C6 halocycloalkyl, CN, SCN, NO2, $N(R^5)R^6$, OR^5 , $C(O)R^5$, $C(O)OR^5$, $C(O)N(R^5)R^6$, SR^5 , $S(O)R^5$, $S(O)_2R^5$. $S(O)_2N(R^5)R^6$ and C_1 - C_6 alkyl substituted with 1 or 2 groups independently selected from NO2, CN, C1-C3 alkylthio, C1-C3 alkoxy. 25 C_1 - C_3 haloalkoxy, C_2 - C_4 alkylearbonyl and C_2 - C_4 alkoxycarbonyl; R3, which is attached to any carbon of the azacyclic ring including the carbon

directly attached to the heterocyclic aromatic ring, is selected from the group H, halogen, C_1 - C_6 alkyl, C_1 - C_6 haloalkyl, C_2 - C_5 alkenyl, C_2 - C_6

BEST AVAILABLE COPY

53

helozikenyl, Ca-C6 alkynyl, Ca-C6 haloalkynyl, Ca-C6 cycloalkyl, Ca-C6 haloeveloulkyl, CN, SCN, NO2, N(R7)R8, OR7, C(O)R7, C(O)OR7, $C(O)N(R^7)R^8$, SR^7 , $S(O)R^7$, $S(O)_2R^7$, $S(O)_2N(R^7)R^8$ and C_1-C_6 alkyl substituted with 1 or 2 groups independently selected from the group NO2, CN, C_1 - C_3 alkylthio, C_1 - C_3 alkoxy, C_4 - C_3 haloalkoxy, C_2 - C_4 alkylcarbonyl 5 and Co-Ca alkoxycarbonyl; R4 and R11 are independently selected from the group H, C1-C6 alkyl, C3-C6 cycloalkyl, C1-C6 haloalkyl, C2-C6 alkenyl, C2-C6 alkynyl, N(R9)R10, $C(O)R^9$, $C(O)OR^9$, $C(O)N(R^9)R^{10}$, SR^9 , $S(O)R^9$, $S(O)_2R^9$. S.O.:-N(R⁹)R¹⁰, benzyl and CH(CH₃)Ph; provided when any of R¹, R², R³ 10 or R4 is $S(O)R^5$, $S(O)_2R^5$, $S(O)R^7$, $S(O)_2R^7$, $S(O)R^9$, or $S(O)_2R^9$ then R5, R and R9 are other than H; R5, R6, R7, R8, R9 and R10 are independently selected the group H, C1-C6 alkyl, C₁-C₅ haloalkyl, C₃-C₆ cycloalkyl, phenyl optionally substituted with 1 or 2 substituents independently selected from W, and benzyl optionally 15 substituted with 1 or 2 substitutents independently selected from W; W is selected from the group halogen, NO2, CN, C1-C3 alkyl, C1-C3 haloalkyl, C_1 - C_3 alkylthio, C_1 - C_3 alkoxy, C_1 - C_3 haloalkoxy, C_2 - C_4 alkylcarbonyl and C₂-C₂ alkoxycarbonyl; m and a are independently 0, 1 or 2; 20 p is 1 c: 2; and q is 1, 2 or 3. A method according to Claim 5 wherein: 6. R^{\pm} is selected from the group H, halogen and C_1 - C_2 alkyl; 25 R2 is selected from the group H and Cl; R3 is selected from the group H, halogen, C₁-C₆ alkyl and OR⁷; R5 is selected from the group H and C1-C4 alkyl; and m and n are independently 0 or 1. 30 A method according to Claim 6 wherein: V is N: $X \text{ is } -C(R^{\frac{1}{2}})=C(R^{\frac{1}{2}})-1$ Y and Z are -C(R1)-; and 35 Q1 is Q-6.

> A method according to Claim 6 wherein: V is N;

X is S: Y and Z are $-C(R^T)$ -; and Q^T is Q-6.

- 5 9. A method according to Claim 6 wherein:
 the ring contains two N and one O or S; and
 Q1 is Q-6.
- 10. An arthropodicidal composition comprising a carrier and an arthropodicidally effective amount of a compound of the formula:

$$V - Z$$
 $V - Z$
 $V - Z$
 $V - Z$
 Q

1

wherein:

15 Q1 is selected from the group

$$(R^{3})_{p} = (CH_{2})_{n}$$

$$(R^{3})_{p} = (CH_{2})_{q}$$

$$(CH_{2})_{q}$$

where the broken line represents an optional chemical bond;

V. X. Y and Z of the ring are each independently selected from the group O, S. N. -C(R¹)-, -C(R¹)=C(R²)-, -C(R¹)=N- and -N(R¹¹)-; provided that (i) no more than one of V, X, Y or Z is -C(R¹)=C(R²)-, -C(R¹)=N-, -N(R¹¹)-. O or S. (ii) at least one of V, X, Y or Z is N, (iii) when the ring is a 5-membered ring containing two N and one O or S, and Q is Q-6, then the ring is attached to the 2-position of Q¹ and (iv) when the ring is a 5-membered ring containing two N and one O or S, then Q is other than Q-5;
R¹ and R² are independently selected from the group W belongs G, C, C, W, the selected ring containing two N and one O or S, then Q is other than Q-5;

R1 and R2 are independently selected from the group H, halogen, C_1 - C_6 alkyl, C_1 - C_5 haloalkyl, C_2 - C_6 alkenyl, C_2 - C_6 haloalkenyl, C_2 - C_6 alkynyl, C_3 - C_6 haloalkyl, C_3 - C_6 haloalkyl, C_3 - C_6 haloayeloalkyl, C_3 - C_6

Elling.

	$N(R^{\frac{1}{2}}, R^{n}, OR^{\frac{1}{2}}, C(O)R^{\frac{1}{2}}, C(O)OR^{\frac{1}{2}}, C(O)N(R^{\frac{1}{2}})R^{n}, SR^{\frac{1}{2}}, S(O)R^{\frac{1}{2}},	-----	--
	SiOi2NiR5iR6 and C1-C6 alkyl substituted with 1 or 2 groups		
	independentity selected from NO2, CN, C1-C3 alkylthio, C1-C3 alkoxy.		
	C_2 : haloutkoxy, C_2 - C_4 alkyleurbonyl and C_2 - C_4 alkoxyeurbonyl:		
5	R4, which is attached to any carbon of the azacyclic ring including the carbon		
	directly attached to the heterocyclic aromatic ring, is selected from the		
	group H. halogen, C ₁ -C ₅ alkyl, C ₁ -C ₅ haloalkyl, C ₂ -C ₆ alkenyl, C ₂ -C ₅		
	haloalkenyl, C ₂ -C ₆ alkynyl, C ₂ -C ₆ haloalkynyl, C ₃ -C ₆ eyeloalkyl, C ₃ -C ₆		
	halocycloaikyl, CN, SCN, NO ₂ , N(R ⁷)R ⁸ , OR (C(O)R ⁷ , C(O)OR ⁷ ,		
10	$C(O,N;R^7;R^8,SR^7,S(O)R^7,S(O)_2R^7,S(O)_2N(R^7)R^8$ and C_1 - C_6 alkyl		
	substituted with 1 or 2 groups independently selected from NO2, CN.		
	C1-C3 alky ithio, C1-C3 alkoxy, C1-C3 haloalkoxy, C2-C4 alkylearbonyl and		
	CCz alkoxycurbonyli		
	R4 and R14 are independently selected from the group H, C1-C6 alkyl, C2-C6		
15	cycloalkyl, C ₁ -C ₅ haloalkyl, C ₂ -C ₆ alkenyl, C ₂ -C ₅ alkynyl, N(R ⁹)R ¹⁰ ,		
	C(O)R", C(O)OR", C(O)N(R9)R10, SR9, S(O)R9, S(O)2R9.		
	S(O) ₂ N(R ⁴)R ⁴ , benzyl and CH(CH ₄ -Pit; provided when any of R ⁴ , R ² , R ³		
	or R^4 is $S(O)R^5$, $S(O)_2R^5$, $S(O)R^7$, $S(O)_3R^7$, $S(O)R^9$, or $S(O)_2R^9$ then R^5 .		
	R ⁷ and R ⁹ are other than H;		
-20	R^5 , R^6 , R^7 , R^8 , R^9 and R^{10} are independently selected the group H, C_1 - C_6 alkyl,		
	C_1 - C_6 haloalkyl, C_5 - C_6 cycloalkyl, phenyl optionally substituted with 1 or 2		
	substituents independently selected from W, and benzyl optionally.		
	substituted with 1 or 2 substitutents independently selected from Wi		
	Was selected from the group halogen, NO ₂ , CN, C ₁ -C ₃ alkyl, C ₁ -C ₃ haloalkyl,		
25	C_1 - C_3 alkylihio, C_4 - C_3 alkoxy, C_4 - C_3 haloalkoxy, C_2 - C_4 alkylearbonyl and		
	C ₂ -C ₄ alkoxycarbonyl;		
	in and a are independently 0. 1 or 2;		

m and n are independently 0. 1 or 2 p is 1 or 2; and q is 1, 2 or 3

INTERNATIONAL SEARCH REPORT

International application ha. PCT/US 94/08404

A. CLASSIFICATION OF SUBJECT MATTER
IPC 6 C07D487/08 C07D453/06
C07D453/02 //(C07D487/ A01N43/82 A01N43/90 //(C070487/08,209:00,209:00)

C07D471/08

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Miramum documentarion searched (classification system (dilowed by classification symbols) IPC 6 CO7O A01N

Documentation searched other than ministrum excurrentation to the extent that such documents are included in the fields searched

Electronic data base consulted curing the international search (name of data base and, where practical, search terms used)

C	Grazon of document, with the exercise, where appropriate, of the relevant passages	Relevant to claim No.
CTREMA	CII.S. J. SCCIRIL STOP	
Х,Р	WO,A,93 14635 (COWELANCO) 5 August 1993 cited in the application see claim 1	5
X	CHEMICAL ABSTRACTS, vol. 62, no. 8, 1965, Columbus, Ohio, US; abstract no. 9101c, A.S. SADYKOV ET AL. 'Syntheses based on anabasine. XIX. Synthesis of 7-methylquinuclidine and alpha-(7-methylquinuclidyl)-beta-pyridine' see abstract & ZH. OBSHCH. KHIM. 34(12), 4104-7 (1964)	1-3

Further documents are lined in the continuation of box C.	Patent family members are listed in armos.
* Special categories of cited documents: *A* document selling the general rate of the art which is not connidered to be of paracular relevance. *S* earlier document but published on or after the international filing date. *L* document which may throw document one on or after the categories of entire to creation or processing date of another categories of other special research in operation. *O* document referring to an oral discussion, use, excition or other means. *P* document ordering do an oral discussion of filing date but later than the property date diamets.	The later document published after the international filing date or priority date and not in conflict with the application but died to understand the principle or theory underlying the invention. "X" document of particular minimates; the diamed invention cannot be considered movel or cannot be considered to involve an inventive step when the document is taken alone. "Y" document of particular relevances the diamed invention cannot be considered to involve an inventive step when the document is combined with one or more other such discussions and the art. "S" document incombined only or your family.
Date of the sexual complexes of the international watch	Date of mailing of the international search report
8 November 1994	1 7. 11. 94
Name and making address of the IBA European Patent Office, P.B. 31, 3 Patentiases 1	Authorized afficat
NL - 2210 HV Reports Tel. (= 31.70) 140 max. Tel. 31 431 494 ml. 914 (= 31.70) 340-3314	Alfaro Faus, I

Form POT ISA 312 (seeme shoul) (July 1972)

INTERNATIONAL SEARCH REPORT

... formation on patent family members

International application No. PCT/US 94/08404

WO-A-9314636	Patent document clied in search report	Publication date	Patent family member(s)		Publication date	
Et V 0311100 15 01 1	WO-A-9314636	05-08-93	-8-UA -8-UA	651516 3239693	21-07-94 01-09-93	

-CT NAMABLE COPY

Form FCT ISA III cratery carrier anneal (Act) 1881:

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US 94/08404

Box I Observations where extrain daims w	ere found unsearchable (Continuation of item 1 of first sheet)
This international search report has not been enai-	elished in respect of certain cisies under Article 17(2)(x) for the following ressons:
Claims Nosa because they relate to subject matter not	required to be searched by this Authority, namely:
2. Cisi=s N≃::	required to be searched by this Authority, namely: Autoral application that do not comply with the prescribed requirements to such all search can be carried out, specifically:
The scope of claim 5 is the POT (conciseness of the PPO Part B. Chapt.	so broadly formulated that on grounds of Art. 6 of claims) and of the Guidelines for Examination in III, 2.2 (economic reasons) the search has been sclosed in the description.
Cisims Note: because they are dependent cours and a	er not drafted in apportance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention	en is leaking (Continuetion of item 2 of first sheet)
This International Sourching Authority found ma	Caple avenuons in this international application, as follows:
	·
	·
1. As all recovered additional search feet we search this claums.	ere amely paid by the applicant, this international resuch report covers all
2. As all marchable dams only be some of any accurant fee.	nes ಕಾಯಿಂದ eñost justifying en entimonal fee, this Authority ಮೆರೆ not ಹಾಗೆಯ payment
3 As only name of the requires stations worts only those distinct for which for	ri seruch feer water omely brid på nur robboard pprintentationer terrep ubbott
4. No resum waters artifes we retried to the averages for means	m turnily puld by the apprount Consequently, this international searth report is shed in the course; it is covered by claims Nosi:
Rouse, on Probat	The administrative was some amomparied by the applicant's protest. No protest amomparies the payment of administrative feet.

Form FCT ISA 200 management of Processes (1977, 2019, 1992).